SURGICAL STAPLING INSTRUMENT WITH IMPROVED FIRING TRIGGER ARRANGEMENT

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Appl. No.: 12/855,351
Filed: Aug. 12, 2010

Related U.S. Application Data
Continuation-in-part of application No. 12/725,993, filed on Mar. 17, 2010, which is a continuation-in-part of application No. 12/234,149, filed on Sep. 19, 2008, Continuation-in-part of application No. 12/622,099, filed on Nov. 19, 2009, Continuation-in-part of application No. 12/843,436, filed on Jul. 26, 2010, which is a continuation of application No. 12/030,424, filed on Feb. 13, 2008, now Pat. No. 7,766,209.

Publication Classification
Int. Cl. A61B 17/068 (2006.01)
U.S. Cl. 227/178.1

ABSTRACT
A surgical stapling instrument including an actuator knob which can be moved from one side of the stapling instrument to another side in order to reposition the actuator knob without having to reposition the stapling instrument within a surgical site. A stapling instrument can include a pusher bar, a housing having a first side and a second side, and an actuator knob rotatably mounted to the pusher bar wherein the actuator knob can be configured to be rotated between a first position in which the actuator knob can be moved along the first side of the housing and a second position where the actuator knob can be moved along a second side of the housing. Alternatively, a surgical stapling instrument can comprise one or more actuator knobs which can be operably engaged and disengaged with a pusher bar in order to selectively utilize the actuator knobs.
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CROSS-REFERENCE TO RELATED APPLICATIONS


BACKGROUND

[0002] i. Technical Field
[0003] The present invention relates to stapling instruments and, in various embodiments, to a surgical stapling instrument for producing one or more rows of staples.
[0004] ii. Background of the Related Art
[0005] In recent years, there has been an increasing tendency for surgeons to use stapling instruments to suture body tissues such as a lung, an esophagus, a stomach, a duodenum and/or other organs in the intestinal tract. The use of an appropriate stapling instrument in many instances may perform a better job in less time and simplify previously difficult surgical procedures such as gastrointestinal anastomoses. Previous linear two and four row cutting staplers comprised cartridge-less instruments into which staples were individually hand-loaded. Other previous devices have included a presterilized disposable staple loading unit and a cutting member which could be utilized for dividing the tissue and forming the rows of staples simultaneously. An example of such a surgical stapler is disclosed in U.S. Pat. No. 3,499,591, entitled INSTRUMENT FOR PLACING LATERAL GASTROINTESTINAL ANASTOMOSES, which issued on Mar. 10, 1970, the entire disclosure of which is hereby incorporated by reference herein.
[0006] A stapling instrument can include a pair of cooperating elongate jaw members, wherein each jaw member can be adapted to be inserted into an internal, tubular body organ to be anastomosed. In various embodiments, one of the jaw members can support a staple cartridge with at least two laterally spaced rows of staples, and the other jaw member can support an anvil with staple-forming pockets aligned with the rows of staples in the staple cartridge. Generally, the stapling instrument can further include a pusher bar and knife blade which are slideable relative to the jaw members to sequentially eject staples from the staple cartridge via camming surfaces on the pusher bar. In at least one embodiment, the camming surfaces can be configured to activate a plurality of staple drivers carried by the cartridge and associated with the individual staples to push the staples against the anvil and form laterally spaced rows of deformed staples in the tissue gripped between the jaw members. In typical stapling instruments, however, the anvil is immovable relative to the staple cartridge once the jaw members have been assembled together and the formed height of the staples cannot be adjusted. In at least one embodiment, the knife blade can trail the pusher bar and cut the tissue along a line between the staple rows. Examples of such stapling instruments are disclosed in U.S. Pat. No. 4,429,695, entitled SURGICAL INSTRUMENTS, which issued on Feb. 7, 1984, the entire disclosure of which is hereby incorporated by reference herein.

SUMMARY

[0007] In at least one form, a surgical stapling instrument can comprise a first side, a second side, a longitudinal axis, a first housing member, and a second housing member. The first housing member can comprise a staple cartridge attachment portion configured to operably support a staple cartridge, and a first lock rail extending along the longitudinal axis. The second housing member can comprise a jaw member configured to operably support an anvil configured to deform staples deployed from the staple cartridge, and a second lock rail extending along the longitudinal axis. The surgical stapling instrument can further comprise a pusher bar configured to move a staple driver relative to the staple cartridge attachment portion and the jaw member, and an actuator configured to receive a force thereto, wherein the actuator is configured to be moved between a first position and a second position, wherein the actuator is configured to be moved along the first side when the actuator is in the first position, wherein the actuator is configured to be moved along the second side when the actuator is in the second position, and wherein the actuator further comprises at least one lock rail receiver configured to receive the first lock rail and the second lock rail.

[0008] In at least one form, a surgical stapling instrument can comprise a first side, a second side, a first housing member, and a second housing member. The first housing member can comprise a first proximal end portion, a first distal end portion, a staple cartridge attachment portion configured to operably support a staple cartridge, and a first longitudinal lock portion extending from the first proximal end portion toward the first distal end portion. The second housing member can comprise a second proximal end portion, a second distal end portion, a jaw member configured to operably support an anvil configured to deform staples deployed from the staple cartridge, and a second longitudinal lock portion extending from the second proximal end portion toward the second distal end portion. The surgical stapling instrument can further comprise a drive member configured to move a staple driver relative to the staple cartridge attachment portion and the jaw member, and an actuator configured to receive a force thereto, wherein the actuator is configured to be moved between a first position and a second position, wherein the
actuator is configured to be moved along the first side when the actuator is in the first position, wherein the actuator is configured to be moved along the second side when the actuator is in the second position, and wherein the actuator further comprises at least one actuator lock portion configured to be aligned with the first longitudinal lock portion and the second longitudinal lock portion.  

In at least one form, a surgical stapling instrument can comprise a first side, a second side, a longitudinal axis, a first housing member, and a second housing member. The first housing member can comprise a staple cartridge attachment portion configured to operably support a staple cartridge, and a first lock rail extending along the longitudinal axis. The second housing member can comprise a jaw member configured to operably support an anvil configured to deform staples deployed from the staple cartridge, and a second lock rail extending along the longitudinal axis. The surgical stapling instrument can further comprise a pusher bar configured to move a staple drive relative to the staple cartridge attachment portion and the jaw member, and an actuator configured to receive a force thereto, wherein the actuator is configured to be moved between a first position and a second position, wherein the actuator is configured to be moved along the first side when said actuator is in the first position, wherein the actuator is configured to be moved along the second side when the actuator is in the second position, and wherein the actuator further comprises capturing means for capturing the first lock rail and the second lock rail.

BRIEF DESCRIPTION OF DRAWINGS

The above-mentioned and other features and advantages of this invention, and the manner of attaining them, will become more apparent and the invention itself will be better understood by reference to the following description of embodiments of the invention taken in conjunction with the accompanying drawings, wherein:

FIG. 1 is a perspective view of a linear anastomotic stapling instrument;

FIG. 2 is a side elevational view showing the anastomotic stapling instrument of FIG. 1 partially disassembled with its upper anvil carrying jaw member detached from its lower staple cartridge carrying jaw member;

FIG. 3 is a side elevational view showing the anastomotic stapling instrument of FIG. 1 in its assembled configuration;

FIG. 4 is a cross-sectional view of the anastomotic stapling instrument of FIG. 1 showing a cam mechanism for urging the rear portions of the upper and lower jaw members apart;

FIG. 5 is a bottom view of the anvil carrying jaw member of the anastomotic stapling instrument of FIG. 1;

FIG. 6 is a top view of the staple cartridge carrying jaw member of the anastomotic stapling instrument of FIG. 1;

FIG. 7 is a bottom view of the anastomotic stapling instrument of FIG. 1;

FIG. 8 is a front end view of the anastomotic stapling instrument of FIG. 1;

FIG. 9 is a rear end view of the anastomotic stapling instrument of FIG. 1;

FIG. 10 is a perspective view of a pusher bar and knife blade assembly of the anastomotic stapling instrument of FIG. 1;

FIG. 11 is a perspective view of a pusher block and an actuator knob which are components of the pusher bar and knife blade assembly of FIG. 10;

FIG. 12 is a partial cross-sectional view of the rear portion of the anastomotic stapling instrument of FIG. 1 illustrating the cam mechanism in its inoperative position;

FIG. 13 is a partial cross-sectional view of the rear portion of the anastomotic stapling instrument of FIG. 1 illustrating the cam mechanism in its operative position;

FIG. 14 is a side view of the staple cartridge of the anastomotic stapling instrument of FIG. 1;

FIG. 15 is a top view of the staple cartridge of the anastomotic stapling instrument of FIG. 1;

FIG. 16 is a bottom view of the staple cartridge of the anastomotic stapling instrument of FIG. 1;

FIG. 17 is a partial cross-sectional view of the anvil and staple cartridge carrying jaw members of FIGS. 5 and 6 illustrating the operation of the pusher bar and knife blade assembly of FIG. 10;

FIG. 18 is a cross-sectional view of the anastomotic stapling instrument of FIG. 1 taken along line 18-18 in FIG. 4;

FIG. 19 is a cross-sectional view of the anastomotic stapling instrument of FIG. 1 taken along line 19-19 in FIG. 4;

FIG. 20 is a detail view of a portion of the anvil and staple cartridge shown in FIG. 18;

FIG. 21 is a perspective view of a stapling instrument in accordance with one non-limiting embodiment of the present invention;

FIG. 22 is a perspective view of the stapling instrument of FIG. 21 illustrating a first actuator knob in an extended position;

FIG. 23 is a perspective view of the stapling instrument of FIG. 21 illustrating the extended actuator knob of FIG. 22 after it has been advanced distally;

FIG. 24 is an exploded view of a clutch mechanism for operably engaging one or more actuator knobs with a pusher bar of the stapling instrument of FIG. 21;

FIG. 25 is a perspective view of a guide member of the clutch mechanism of FIG. 24;

FIG. 26 is a perspective view of an actuator knob of the stapling instrument of FIG. 21;

FIG. 27 is another perspective view of the clutch mechanism of FIG. 24;

FIG. 28 is a perspective view of the stapling instrument of FIG. 21 illustrating the first actuator knob in a retracted position and a second actuator knob in an extended position;

FIG. 29 is a partial exploded view of a stapling instrument in accordance with one non-limiting embodiment of the present invention;

FIG. 30 is a partial perspective view of the stapling instrument of FIG. 29 illustrating an actuator knob after it has been advanced distally along a first side of the stapling instrument;

FIG. 31 is a partial perspective view of the stapling instrument of FIG. 29 illustrating the actuator knob of FIG. 30 being rotated between a first position and a second position;

FIG. 32 is a partial perspective view of the stapling instrument of FIG. 29 illustrating the actuator knob of FIG. 30 after it has been advanced distally along a second side of the stapling instrument;
FIG. 33 is an exploded view of a pusher bar assembly of the stapling instrument of FIG. 29 configured to allow the actuator knob of FIG. 30 to be rotated between its first and second positions;

FIG. 34 is a perspective view of a surgical stapling instrument in accordance with at least one embodiment of the present invention;

FIG. 35 is an exploded perspective view of the surgical stapling instrument of FIG. 34;

FIG. 36 is an exploded elevational view of the surgical stapling instrument of FIG. 34;

FIG. 37 is a partial cross-sectional view of the surgical stapling instrument of FIG. 34 illustrating first and second portions being assembled together;

FIG. 38 is a partial cross-sectional view of the surgical stapling instrument of FIG. 34 illustrating the proximal end of the first portion of FIG. 37 being locked to the proximal end of the second portion of FIG. 37 and illustrating the second portion being rotated toward the first portion;

FIG. 39 is a partial cross-sectional view of the surgical stapling instrument of FIG. 34 illustrating a latch rotatably mounted to the first portion, wherein the latch is engaged with the second portion and wherein the latch has been rotated into a partially-closed position;

FIG. 40 is a partial cross-sectional view of the surgical stapling instrument of FIG. 34 illustrating the latch of FIG. 39 in a closed position;

FIG. 41 is a perspective view of a staple cartridge assembly of the surgical stapling instrument of FIG. 34;

FIG. 42 is an exploded view of the staple cartridge assembly of FIG. 41;

FIG. 43 is a cross-sectional view of the staple cartridge assembly of FIG. 41 taken along line 43-43 in FIG. 42;

FIG. 44 is an exploded view of a staple sled and cutting member assembly of the staple cartridge assembly of FIG. 41;

FIG. 45 is a perspective view of the staple sled and cutting member assembly of FIG. 44;

FIG. 46 is a perspective view of the surgical stapling instrument of FIG. 34 illustrating a firing actuator moved distally along a first side of the surgical stapling instrument;

FIG. 47 is a perspective view of the surgical stapling instrument of FIG. 34 illustrating the firing actuator of FIG. 46 moved distally along a second side of the surgical stapling instrument;

FIG. 48 is a cross-sectional view of a surgical stapling instrument in accordance with at least one alternative embodiment of the present invention illustrating a latch in a partially-closed position and a locking mechanism engaged with a firing actuator;

FIG. 49 is a cross-sectional view of the surgical stapling instrument of FIG. 48 wherein the latch has been moved into a closed position and has disengaged the locking mechanism from the firing actuator;

FIG. 50 is a perspective view of an anvil assembly of the surgical stapling instrument of FIG. 34;

FIG. 51 is an exploded perspective view of the anvil assembly of FIG. 50;

FIG. 52 is another exploded perspective view of the anvil assembly of FIG. 50;

FIG. 53 is an exploded cross-sectional elevational view of the anvil assembly of FIG. 50;

FIG. 54 is a cross-sectional assembly view of the anvil assembly of FIG. 50 illustrating an anvil adjustment member in a first position;

FIG. 55 is a cross-sectional assembly view of the anvil assembly of FIG. 50 illustrating the anvil adjustment member of FIG. 54 in a second position;

FIG. 56 is a cross-sectional assembly view of the anvil assembly of FIG. 50 illustrating the anvil adjustment member of FIG. 54 in a third position;

FIG. 57 is a perspective view of a surgical stapling instrument in accordance with at least one alternative embodiment of the present invention;

FIG. 58 is a cross-sectional view of the surgical stapling instrument of FIG. 57 taken along line 58-58 in FIG. 57;

FIG. 59 is a partial exploded view of the proximal end of the surgical stapling instrument of FIG. 57 including a detent mechanism for releasably holding a rotatable anvil adjustment member in position;

FIG. 60 is a perspective view of the surgical stapling instrument of FIG. 57 with some components removed and others shown in cross-section;

FIG. 61 is an exploded view of portions of the surgical stapling instrument of FIG. 57 illustrating a rotatable anvil adjustment member in a first orientation;

FIG. 62 is a perspective view of the rotatable anvil adjustment member of FIG. 61;

FIG. 63 is an end view of the surgical stapling instrument of FIG. 57 with some components removed and others shown in dashed lines illustrating the rotatable anvil adjustment member in the first orientation of FIG. 61;

FIG. 64 is a cross-sectional end view of the surgical stapling instrument of FIG. 57 taken along line 64-64 in FIG. 57;

FIG. 65 is an end view of the surgical stapling instrument of FIG. 57 illustrating the rotatable anvil adjustment member of FIG. 61 rotated in a first direction into a second orientation;

FIG. 66 is a cross-sectional end view of the surgical stapling instrument of FIG. 57 illustrating the anvil adjustment member in the second orientation of FIG. 65;

FIG. 67 is an end view of the surgical stapling instrument of FIG. 57 illustrating the rotatable anvil adjustment member of FIG. 61 rotated in a second direction into a third orientation;

FIG. 68 is a cross-sectional end view of the surgical stapling instrument of FIG. 57 illustrating the anvil adjustment member in the third orientation of FIG. 67;

FIG. 69 is a perspective view of an actuator for rotating the anvil adjustment member of FIG. 61;

FIG. 70 is a partial cross-sectional view of a surgical stapling instrument including a spring configured to bias the distal end of a first handle portion away from the distal end of a second handle portion when the stapling instrument is in a partially-closed configuration;

FIG. 71 is a similar perspective view of the surgical stapling instrument of FIG. 34 to that of FIG. 50;

FIG. 72 is a detail view of a latch projection extending from an anvil of a surgical stapling instrument in accordance with at least one alternative embodiment of the present invention;

FIG. 73 is a diagram illustrating the latch projection of FIG. 72 and a latch configured to engage the latch projection and move the latch projection into a latch recess;
[0084] FIG. 74 is an elevational view of the latch projection of FIG. 72;  
[0085] FIG. 75 is a perspective view of a staple pocket in accordance with at least one embodiment of the present invention;  
[0086] FIG. 76 is a top view of the staple pocket of FIG. 75;  
[0087] FIG. 77 is a cross-sectional view of the staple pocket of FIG. 75 taken along line 77-77 in FIG. 76;  
[0088] FIG. 78 is a cross-sectional view of the staple pocket of FIG. 75 taken along line 78-78 in FIG. 76;  
[0089] FIG. 79 is another top view of the staple pocket of FIG. 75;  
[0090] FIG. 80 is a cross-sectional view of the staple pocket of FIG. 75 taken along line 80-80 in FIG. 79;  
[0091] FIG. 81 is a cross-sectional view of the staple pocket of FIG. 75 taken along line 81-81 in FIG. 79;  
[0092] FIG. 82 is an elevation view of a surgical staple in an undeformed shape;  
[0093] FIG. 83 is an elevation view of the surgical staple of FIG. 82 in a deformed shape in accordance with at least one embodiment of the present invention;  
[0094] FIG. 84 is a side view of the surgical staple of FIG. 82 in the deformed shape of FIG. 83;  
[0095] FIG. 85 is a plan view of the surgical staple of FIG. 82 in the deformed shape of FIG. 83;  
[0096] FIG. 85A is another plan view of the surgical staple of FIG. 82 in the deformed shape of FIG. 83;  
[0097] FIG. 86 is an elevation view of a surgical staple in an undeformed shape;  
[0098] FIG. 87 is a bottom view of the surgical staple of FIG. 86 in an undeformed shape;  
[0099] FIG. 88 is a bottom view of the surgical staple of FIG. 86 in a deformed shape in accordance with at least one embodiment of the present invention;  
[0100] FIG. 89 is a partial cross-sectional view of the surgical staple of FIG. 86;  
[0101] FIG. 90 is an elevation view of a surgical staple in a deformed shape in accordance with at least one embodiment of the present invention;  
[0102] FIG. 91 is an elevation view of a surgical staple in a deformed shape;  
[0103] FIG. 92 is an exploded perspective view of the surgical stapling instrument of FIG. 34;  
[0104] FIG. 93 is an exploded elevation view of the surgical stapling instrument of FIG. 34;  
[0105] FIG. 94 is a partial cross-sectional view of the surgical stapling instrument of FIG. 34 illustrating a latch rotatably mounted to the first portion, wherein the latch is engaged with the second portion and wherein the latch has been rotated into a partially-closed position;  
[0106] FIG. 95 is a perspective view of a staple cartridge assembly of the surgical stapling instrument of FIG. 34;  
[0107] FIG. 96 is an exploded view of the staple cartridge assembly of FIG. 95;  
[0108] FIG. 97 is a cross-sectional view of the staple cartridge assembly of FIG. 95;  
[0109] FIG. 98 is an exploded view of a staple sled and cutting member assembly of the staple cartridge assembly of FIG. 95;  
[0110] FIG. 99 is a perspective view of the staple sled and cutting member assembly of FIG. 98;  
[0111] FIG. 100 is a detail view of a distal end of a drive bar configured to be operably connected to the staple sled and cutting assembly of FIG. 98, wherein the drive bar distal end is illustrated in a proximal position in solid lines a second, or distal, position in phantom lines;  
[0112] FIG. 101 is a partial bottom view of the staple cartridge assembly of FIG. 95;  
[0113] FIG. 102 is a cross-sectional view of a staple cartridge assembly in accordance with an alternative embodiment;  
[0114] FIG. 103 is a perspective view of a surgical stapling instrument comprising a firing actuator in a partially-advanced position;  
[0115] FIG. 104 is a cross-sectional view of the surgical stapling instrument of FIG. 103 illustrating the firing actuator in the partially-advanced position;  
[0116] FIG. 105 is a cross-sectional view of the surgical stapling instrument of FIG. 103 illustrating the firing actuator being returned toward an unfired position;  
[0117] FIG. 106 is a top view of the surgical stapling instrument of FIG. 103 illustrating the firing actuator being moved distally;  
[0118] FIG. 107 is a top view of the surgical stapling instrument of FIG. 107 illustrating the firing actuator being moved proximally;  
[0119] FIG. 108 is another perspective view of the surgical stapling instrument of FIG. 103;  
[0120] FIG. 109 is a cross-sectional view of the proximal end of the surgical stapling instrument of FIG. 103 illustrating the firing actuator in an unfired position;  
[0121] FIG. 110 is a cross-sectional view of the proximal end of the surgical stapling instrument of FIG. 103 illustrating the firing actuator rotated to a first side of the surgical stapling instrument housing; and  
[0122] FIG. 111 is a cross-sectional view of the proximal end of the surgical stapling instrument of FIG. 103 illustrating the firing actuator in a partially-fired position.  
[0123] Corresponding reference characters indicate corresponding parts throughout the several views. The exemplifications set out herein illustrate preferred embodiments of the invention, in one form, and such exemplifications are not to be construed as limiting the scope of the invention in any manner.

DETAILED DESCRIPTION

[0124] Certain exemplary embodiments will now be described to provide an overall understanding of the principles of the structure, function, manufacture, and use of the device and methods disclosed herein. One or more examples of these embodiments are illustrated in the accompanying drawings. Those of ordinary skill in the art will understand that the devices and methods specifically described herein and illustrated in the accompanying drawings are non-limiting exemplary embodiments and that the scope of the various embodiments of the present invention is defined solely by the claims. The features illustrated or described in connection with one exemplary embodiment may be combined with the features of other embodiments. Such modifications and variations are intended to be included within the scope of the present invention.

[0125] The entire disclosures of the following United States patent applications are hereby incorporated by reference herein:

Referring to FIGS. 2 and 6, lower cartridge carrying jaw member 24 can comprise a one-piece elongated channel-shaped frame including a pair of opposed, elongated side walls 52 connected by a bottom wall 53. Along the rearward portion of lower jaw member 24, a pair of spaced, elongated upstanding side flanges 54 can extend upward from its opposed side walls 52. As shown in FIGS. 5 and 6, the width of lower jaw member 24 between its side flanges 54 can be greater than the width of upper jaw member 22 between its side walls 30 to permit the rear portion of the upper jaw member to be received between side flanges 54 of the lower jaw member when the stapling instrument is assembled for operation. As shown in FIG. 2, each side flange 54 of lower jaw member 24 can include a vertical notch 56 located in alignment with latch pin 36 on upper jaw member 22. When upper jaw member 22 and lower jaw member 24 are assembled, the opposite ends of latch pin 36 can be received in notches 56.

As shown in FIGS. 2 and 6, lower jaw member 24 can support a staple cartridge 60 which is adapted to receive a plurality of surgical staples 61 (FIG. 17) arranged in at least two laterally spaced longitudinal rows. Staple cartridge 60 can be mounted at the front portion of lower jaw member 24 between its side walls 52. Staple cartridge 60 can be divided longitudinally by a central, elongated slot 62 (FIG. 6) which extends from the proximal end of the cartridge toward its distal end. In various embodiments, a plurality of staple openings 64 formed in staple cartridge 60 can be arranged in two pairs of laterally spaced rows, with each pair of rows disposed on opposite sides of central longitudinal slot 62. A plurality of surgical staples 61 (FIG. 17) can be mounted within openings 64 of cartridge 60. As shown in FIG. 6, the staple openings 64 in adjacent rows can be staggered to provide more effective stapling of the tissue when the instrument is operated. Referring to FIGS. 15 and 16, staple cartridge 60 can include a pair of longitudinal slots 66 located on opposite sides of elongated central slot 62 and disposed between the staggered rows of openings 64 on each side of the central slot. Each longitudinal slot 66 can extend from the proximal end of cartridge 60 towards its distal end.

As shown in FIG. 17, a plurality of staple drivers 65 can be slidably mounted in staple openings 64 for actuating the staples 61 which are loaded into staple cartridge 60. Referring to FIG. 6, each staple driver 65 can be designed to simultaneously actuate two staples 61 located in the adjacent rows provided in staple cartridge 60. Thus, in various embodiments, a first set of staple drivers 65 can be provided for actuating the staples 61 in the staggered rows located on one
side of central longitudinal slot 62, and a second set of staple drivers 65 can be provided for actuating the staples 61 in the pair of adjacent rows located on the other side of central longitudinal slot 62.

[0143] As shown in FIGS. 2 and 3, similar to the above, the front or distal end of staple cartridge 60 can include a tapered tip 68 to facilitate the insertion of lower jaw member 24 into a hollow, tubular body organ, for example. Immediately behind its tapered tip 68, staple cartridge 60 can be provided with a pair of rearwardly extending protrusions 70 (one shown in FIG. 14) which can be received in corresponding notches provided in side walls 52 of lower jaw member 24. At the rear of staple cartridge 60, a pair of depending arms 72 can extend downwardly from the cartridge. Each arm 72 can be notched to provide a side opening 74. When cartridge 60 is assembled on lower jaw member 24, its protrusions 70 can be received in corresponding notches provided at the front ends of side walls 52 and its depending arms 72 extend downwardly through an opening 76 (FIG. 4) formed in bottom wall 53 of jaw member 24. Lower jaw member 24 can include a pair of depending ears 78 (FIG. 18) extending downwardly from its side walls 52 on opposite sides of opening 76. A pivot pin 80 can extend through holes formed in depending ears 78 of lower jaw member 24 and through side openings 74 of depending arms 72 on staple cartridge 60 to fasten the staple cartridge to the lower jaw member.

[0144] Referring to FIG. 2, the stapling instrument 20 can include a latching mechanism 90, for latching upper jaw member 22 and lower jaw member 24 together at an intermediate position along the jaw members. In various embodiments, jaw members 22 and 24 can be latched together at a position adjacent to the proximal ends of anvil 40 and staple cartridge 60. In at least one embodiment, latching mechanism 90 can comprise a latch arm 92 (FIG. 2) pivotally connected to lower jaw member 24 via pivot pin 80 (FIG. 4). Latch arm 92 can be channel-shaped in configuration and can include a pair of opposed, elongated side walls 94 (FIG. 6) which are spaced apart by a distance sufficient to span side walls 52 of lower jaw member 24. Each side wall 94 of latch arm 92 can include an upwardly and forwardly extending hook member 96 provided with a forwardly facing slot 98 for receiving latch pin 36. A shroud 100 can be mounted on the lower surface of latch arm 92. When latch arm 92 is closed, as shown in FIG. 3, shroud 100 can be aligned with the bottom of lower handle 28 to facilitate the handling and operation of stapling instrument 20 by the surgeon. In various embodiments, shroud 100 can be made of plastic or other lightweight materials, for example, while latch arm 92 can be made of stainless steel, for example. As shown in FIG. 7, shroud 100 can include elongated flanges 102 and 104 extending outwardly from its opposite sides which can serve as fingers to enable latch arm 92 to be pivoted downwardly from its latched to its unlatched position. When latch arm 92 is moved to its closed or latched position, the surfaces of slots 98 of hook members 96 can cooperate with latch pin 36 which can act as an over-center latch to maintain latch arm 92 in its latched position.

[0145] Referring to FIGS. 6 and 10, the preferred embodiment of stapling instrument 20 can include an improved pusher bar and knife blade assembly, generally 110, which can be slidable mounted for longitudinal movement relative to upper and lower jaw members 22 and 24, respectively, for driving staples 61 from staple cartridge 60 into tissue gripped between the jaw members, forming staples 61 against anvil 40, and cutting the tissue along a line between the rows of staples formed in the tissue. Pusher bar and knife blade assembly 110 can include a pusher block 112 (FIG. 6) which can be slidably received within the lower channel-shaped jaw member 24 between its upstanding side flanges 54. As shown in FIG. 11, pusher block 112 can be attached to an actuator knob 114 by a flange 116 which includes a laterally projecting finger 118 provided with a longitudinally extending notch 119 on its top surface. Finger 118 can be snap-fitted into a lateral slot 120 formed in pusher block 112 to locate notch 119 underneath a longitudinal locking bar 121 to secure pusher block 112 and actuator knob 114 together. Flange 116 of actuator knob 114 can extend through and ride along an elongated slot 122 (FIG. 2) formed in side flange 54 of lower jaw member 24.

[0146] The pusher bar and knife blade assembly 110 can include a pair of staple pusher bars 124 (FIG. 10) projecting forwardly from pusher block 112 and slidably received in elongated slots 66 (FIG. 16) of staple cartridge 60. Pusher block 112 can be provided with a pair of vertical slots 126 (FIG. 11) in which pusher bars 124 are secured. As shown in FIG. 10, the front end of each staple pusher bar 124 can be provided with a wedge-shaped tip 128 which defines an inclined cam surface 130 for engaging staple drivers 65 as pusher bars 124 are advanced into staple cartridge 60. As shown in FIG. 21, each staple driver 65 can be provided with a sloped surface 132 oriented at the same angle as cam surface 130 of each staple pusher bar 124 to provide a flat, sliding contact between the surfaces.

[0147] Referring to FIGS. 6 and 10, the pusher bar and knife blade assembly 110 can include a knife block 134 which is slidable mounted for longitudinal movement along lower jaw member 24 between its upstanding side flanges 54. Knife block 134 can include a knife support bar 136 which extends forwardly into central longitudinal slot 62 of staple cartridge 60. An inclined knife blade 138 provided with a beveled cutting edge 140 can be located at the front end of knife support bar 136. The beveled cutting edge of knife blade 138 can be oriented at an angle relative to elongate jaw members 22 and 24 and can be slidably received in central longitudinal slot 62 of staple cartridge 60.

[0148] In various embodiments, knife block 134 can include a pair of longitudinal slots 135 (FIG. 19) extending therethrough which slidably receive staple pusher bars 124 to permit pusher block 112 to slide relative to the knife block. Accordingly, when pusher block 112 is advanced toward staple cartridge 60 by actuator knob 114, staple pusher bars 124 can slide through knife block 134 which remains stationary until the pusher block moves into engagement with the knife block. After knife block 134 is engaged by pusher block 112, the knife block and pusher block can advance simultaneously toward staple cartridge 60. As shown in FIG. 17, knife blade 138 can be advanced through staple cartridge 60 along with staple pusher bars 124, forming staples 61 in the tissue gripped between the jaw members and cutting the tissue between the staple rows. Thereafter, when actuator knob 114 is retracted, pusher block 112 can initially slide staple pusher bars 124 backward through knife block 134 which can remain stationary. Each staple pusher bar 124 can include an offset portion 142 which can move into engagement with knife block 134 after staple pusher bars 124 are withdrawn by a predetermined distance. With offset portions 142 of staple pusher bars 124 engaging knife block 134, pusher block 112 and knife block 134 can be simultaneously
retracted by actuator knob 114 to return pusher bars 124 and knife blade 138 to the start position.

[0149] In accordance with various embodiments of the invention, stapling instrument 20 can be provided with jaw clamping means for applying clamping forces to the jaw members to urge staple cartridge 60 and anvil 40 together during the formation of staples 61. The jaw clamping means can include means for urging the jaw members apart at a position remote from the latching mechanism to resist the forces exerted on staple cartridge 60 and anvil 40 when staples 61 are formed. In at least one embodiment, a cam means can be mounted on one of the jaw members and can be engageable with the other jaw member for moving said jaw members apart at the remote position to urge staple cartridge 60 and anvil 40 together. In various embodiments, a cam member can be pivotally mounted on one of the jaw members at a position remote from the latching mechanism. The cam member can be pivotable from a first inoperative position to a second operative position to move the remote ends of the jaw members apart. The cam member can be operable by pusher block 112 of pusher bar and knife blade assembly 110 to move to its operative position when the pusher block is advanced and to return to its inoperative position when the pusher block is retracted.

[0150] In various embodiments, a cam mechanism, generally 150, can be located adjacent to the rear end of lower jaw member 24, as shown in FIG. 4. Cam mechanism 150 can include a cam member 152 pivotally mounted on a transverse pivot pin 154 extending between upstanding side flanges 54 of lower jaw member 24. Cam member 152 can include a first lower cam surface 156 for engaging top wall 31 of upper jaw member 22 with cam 152 in its first inoperative position (FIG. 12) and a second higher cam surface 158 for engaging the top wall 31 of upper jaw member 22 with cam 152 in its second operative position (FIG. 13). First cam surface 156 can be arranged to maintain upper and lower jaw members substantially parallel with cam 152 in its inoperative position. Second cam surface 158 can be arranged to raise the rear end of upper jaw member 22 by approximately 0.125 inch (3.2 mm), for example, when cam 152 pivots from its inoperative position to its operative position. In addition, upper jaw member 22 can be sufficiently flexible to permit the rear portion of upper jaw member 22 to bend upward away from lower jaw member 24 when cam member 152 is moved from its inoperative position to its operative position.

[0151] As shown in FIG. 4, cam member 152 can include a radially extending notch 160 which divides the cam into a large front finger 162 and a small rear finger 164. Front cam finger 162 can include a flat, rearwardly facing surface 165, and rear cam finger 164 can include a sloped, forwardly facing surface 166. With cam 152 in its inoperative position, front cam finger 162 and rear cam finger 164 can extend downwardly through an elongated slot 168 formed in bottom wall 53 of lower jaw member 24.

[0152] In various embodiments, cam member 152 can be operable by pusher block 112 to move from its inoperative position to its operative position when the pusher block is advanced. As shown in FIG. 11, pusher block 112 can include a pair of rearwardly extending arms 170 which are spaced apart to define a gap 172 therebetween. The rear ends of arms 170 can be connected by a cam actuator pin 174 which extends across gap 172. Referring to FIGS. 4 and 11, with cam member 152 disposed in its inoperative position, front cam finger 162 can extend through gap 172 between arms 170 of pusher block 112, while cam actuator pin 174 can be received in notch 160 between front finger 162 and rear finger 164 of the cam member.

[0153] As shown in FIG. 12, with cam member 152 disposed in its first inoperative position, top wall 31 of upper jaw member 22 can rest on first cam surface 156 of the cam member. With cam member 152 in its inoperative position, top wall 31 of upper jaw member 22 can be substantially parallel to bottom wall 53 of lower jaw member 24. In addition, pusher block 112 can be located in its start position spaced rearwardly from knife block 134. When pusher block 112 is advanced, as indicated by arrow 182 (FIG. 13), cam actuator pin 174 can engage rear surface 165 of front cam finger 162 to rotate cam member 152 in a counter-clockwise direction, as indicated by arrow 184, to pivot the cam member to its second operative position and move its second cam surface 158 into engagement with top wall 31 of upper jaw member 22. With cam member 152 pivoted to its operative position, the top wall 31 of upper jaw member 22 can be bent upwardly, as indicated by arrow 186, away from bottom wall 53 of lower jaw member 24. The cam member can apply forces to upper jaw member 22 and lower jaw member 24 which bend the rear portions of the jaw members apart. As a result of the bending the rear portions of upper jaw member 22 and lower jaw member 24 apart, additional clamping forces can be applied to the front portions of upper jaw member 22 and lower jaw member 24 to clamp anvil 40 and staple cartridge 60 against the tissue gripped between the jaw members. Thus, anvil 40 and staple cartridge 60 can be urged together to resist the forces exerted on the anvil and staple cartridge when pusher bar and knife blade assembly 110 is advanced to form staples 61 and cut the tissue.

[0154] Referring to FIG. 13, when pusher block 112 is retracted after staples 61 are formed, cam actuator pin 174 can engage sloped surface 166 of rear cam finger 164 to pivot cam member 152 in a clockwise direction. As cam actuator pin 174 moves along sloped surface 166 into notch 160, cam member 152 can pivot in a clockwise direction and return to its first inoperative position (FIG. 12) with its first cam surface 156 in engagement with top wall 31 of upper jaw member 22. As a result, the forces exerted on the rear portions of upper jaw member 22 and lower jaw member 24 by cam 152 can be released and top wall 31 of upper jaw member 22 can return to a substantially parallel relationship with bottom wall 53 of lower jaw member 24. Similarly, the clamping forces applied to the front portions of jaw members 22 and 24 can be released to unclamp anvil 40 and staple cartridge 60.

[0155] In various embodiments, stapling instrument 20 can include spacer means mounted on one of the jaw members for maintaining a predetermined gap between staple cartridge 60 and anvil 40 of the stapling instrument. Referring to FIGS. 4 and 6, this spacer means can be embodied as a spacer pin 190 mounted adjacent to the distal end of staple cartridge 60. Spacer pin 190 can extend vertically upward from bottom wall 53 of lower jaw member 24 through staple cartridge 60 and project upwardly from the top of the staple cartridge by a predetermined distance. As shown in FIG. 5, one flange 38 of anvil 40 can include a flange section 192 adjacent to its distal end for engaging spacer pin 190. With the stapling instrument assembled for operation (FIG. 4), spacer pin 190 can engage flange section 192 to maintain a predetermined gap between anvil 40 and staple cartridge 60.

[0156] In the operation of stapling instrument 20, the tissue to be stapled and cut can be initially placed between jaw
members 22 and 24 and clamped by the jaw members. Thus, handles 26 and 28 can be unlatched by pivotal movement of latch arm 92 downward to its unlatched position (FIG. 2). As a result, the opposite ends of latch pin 36 can be disengaged from slots 98 formed in hook members 96 of latching arm 92. Thereafter, upper and lower jaw members 22 and 24 can be separated by disengaging latch pin 36 from slots 56 formed in side flanges 54 of the lower jaw member.

[0157] Next, the tissue to be stapled and cut can be placed on jaw members 22 and 24. For example, as shown in FIG. 17, a piece of tubular, intestinal tissue may be slipped onto the front portion of each jaw member. After the tissue is placed on the jaw member, stapling instrument 20 can be reassembled. The reassembly can be accomplished by aligning latch pin 36 with vertical slots 56 formed in upstanding side flanges 54 of lower jaw member 24. Thereafter, side flanges 54 of lower jaw member 24 can be positioned inside upper handle 26, spanning side walls 30 of upper jaw member 22, while the opposite ends of latch pin 36 can be inserted into vertical slots 56. Finally, latch arm 92 can be pivoted upward to its latched position (FIG. 3) with its cover 100 flush with the bottom of lower handle 28. As a result, hook members 92 can be pivoted over latch pin 36 and slots 98 can receive the opposite ends of the latch pin. Thus, upper jaw member 22 and lower jaw member 24 can be latched together at an intermediate position therealong adjacent to anvil 40 and staple cartridge 60. In addition, spacer pin 190 can engage flange section 192 of anvil 40 through the body tissue to maintain a predetermined gap between anvil 40 and staple cartridge 60.

[0158] After the tissue is clamped between the jaw members, stapling instrument 20 can be fired by actuating actuator knob 114 to actuate the pusher bar and knife blade assembly 110. Initially, in the actuation of cam mechanism 150, pusher block 112 and pusher bars 124 (FIG. 4) can be advanced, while knife block 134 can remain stationary. Since only pusher block 112 and its pusher bars 124 are advanced to actuate cam member 152, the initial force required to operate stapling instrument 20 can be minimized.

[0159] Referring to FIG. 12, during the initial advance of pusher block 112, pusher bars 124 can slide through knife block 134 and the wedge-shaped tips 128 of the pusher bars can begin to advance through slots 66 of staple cartridge 60. As pusher block 112 advances toward knife block 134, its cam actuator pin 174 can engage rear surface 165 of front cam finger 162 to pivot cam 152 counterclockwise, as indicated by arrow 184 of FIG. 13, to move the second cam surface 158 of the cam member into engagement with top wall 31 of upper jaw member 22. Cam member 152 can apply forces to upper jaw member 22 and lower jaw member 24 which bend the rear portions of the jaw members apart. As a result, the rear end of top wall 31 of upper jaw member 22 can be bent upward by approximately 0.125 inch (3.2 mm), for example, relative to the rear end of bottom wall 53 of lower jaw member 24. The bending of the rear ends of jaw members 22 and 24 apart can result in additional clamping forces on the front portions of the jaw members to clamp anvil 40 and staple cartridge 60 against the tissue gripped between the jaw members. These additional clamping forces tend to resist the forces exerted on anvil 40 and staple cartridge 60, while the tissue is cut and staples 61 are formed against anvil 40, to maintain the desired spacing between anvil 40 and staple cartridge 60 to produce formed staples 61 which are substantially uniform in height.

[0160] Referring to FIG. 13, after cam mechanism 150 is actuated, pusher block 112 can subsequently engage knife block 134 to begin the longitudinal movement of knife block 134 toward staple cartridge 60. In various embodiments, the initial spacing between pusher block 112 and knife block 134 can be arranged such that pusher block 112 engages knife block 134 slightly before cam member 152 arrives at its operative position. Alternatively, the initial spacing between pusher block 112 and knife block 134 can be arranged such that pusher block 112 initially engages knife block 134 after the movement of cam member 152 to its operative position is completed. When pusher block 112 engages knife block 134, the advance of knife blade 138 along central longitudinal slots 42 and 62 of anvil 40 and staple cartridge 60, respectively, can be initiated. Thereafter, staple pusher bars 124 and knife blade 138 can be advanced simultaneously to staple and cut the tissue gripped between anvil 40 and staple cartridge 60.

[0161] As pusher block 112 is advanced, staple pusher bars 124 can be moved longitudinally along slots 66 provided in staple cartridge 60. The two wedge-like cam surfaces 130 of staple pusher bars 124 can move through slots 66 into engagement with the sloped surfaces of staple drivers 65 to sequentially drive staples 61 from cartridge 60 and to form staples 61 into B-shaped configuration against anvil flanges 38. The cam surfaces 130 can be located at the same distance from pusher block 112 to simultaneously actuate staple drivers 65 located on opposite sides of central longitudinal slot 62. At the same time, knife block 134 can be advanced to move knife blade 138 through central longitudinal slot 42 of anvil 40 and through central longitudinal slot 62 of staple cartridge 60 to cut the tissue gripped between the jaw members. The additional clamping forces applied to the front portions of upper jaw member 22 and lower jaw member 24 via cam mechanism 150 can tend to resist the forces exerted on anvil 40 and staple cartridge 60 when staples 61 are formed.

[0162] After pusher block 112 is fully advanced to form all of the staples in cartridge 60, the pusher block can be retracted toward its start position by retraction of actuator knob 114. Initially, only pusher block 112 can move backward from staple cartridge 60 because staple pusher bars 124 slide through knife block 134 which remains stationary. When offset portions 142 of staple pusher bars 124 engage the front of knife block 134, the knife block can be moved backward from staple cartridge 60 along with pusher block 112. As a result, staple pusher bars 124 and knife blade 138 can be simultaneously retracted from staple cartridge 60 and anvil 40.

[0163] As pusher block 112 returns toward its start position, cam actuator pin 174 can engage sloped surface 166 of rear cam finger 164 to pivot cam member 152 in a counterclockwise direction toward its operative position. Cam actuator pin 174 can move along sloped surface 166 into slot 160 between cam fingers 162 and 164 to return cam member 152 to its operative position. As a result, second cam surface 158 of cam member 152 can be disengaged from the top wall of upper jaw member 22 and rear end of top wall 31 of upper jaw member 22 and move downwardly into engagement with first cam surface 156. At the same time, front cam finger 162 can pivot downwardly into gap 172 between fingers 170 on pusher block 112, and both cam fingers 162 and 164 can pivot downwardly into slot 168 formed in bottom wall 53 of lower jaw member 24. Thereafter, with cam member 152 in its operative position, latching arm 92 can be pivoted downward, as shown in FIG. 2, to permit upper jaw member 22 and lower jaw member 24 to be disassembled. At this point, the cut and stapled tissue can be removed from the jaw members.
As outlined above, a surgical stapling instrument can include an actuator knob, such as actuator knob 114 (FIG. 1), for example, which can be configured to advance a pusher bar assembly, such as pusher bar assembly 110 (FIG. 10), within a staple cartridge of the surgical stapling instrument. In various embodiments, actuator knob 114 can be configured to be grasped by a surgeon such that the surgeon can apply a force thereto. In various circumstances, actuator knob 114 can come into contact with or about tissue surrounding the surgical site when it is advanced distally. In at least one circumstance, as a result, the surgeon may have to reposition the stapling instrument such that actuator knob 114 can pass by the tissue. In other circumstances, the surgeon may have to force actuator knob 114 by the tissue. In either event, such circumstances may be unsuitable and, as a result, there exists a need for a stapling instrument having an actuator knob which can be manipulated to reduce the possibility that the actuator knob may impinge on the surrounding tissue.

In various embodiments of the present invention, referring to FIG. 21, stapling instrument 220 can include anvil carrying jaw member 222 extending from upper handle 226, staple cartridge carrying jaw member 224 extending from lower handle 228, and actuator knobs 214a and 214b which can be operably engaged with a pusher bar assembly, such as pusher bar assembly 210 as illustrated in FIG. 24, for example. In various embodiments, a staple cartridge can be removably attached to staple cartridge carrying jaw member 224, for example, such that, after the staple cartridge has been expended, it can be replaced with another staple cartridge. In at least one embodiment, pusher bar assembly 210 can include a staple driver integrally-formed with or operably mounted thereto which can be moved through the staple cartridge as outlined above. In at least one other embodiment, the staple cartridge can include a staple driver contained therein which can be engaged with and pushed distally by the pusher bar assembly. In any event, first actuator knob 214a, for example, can be rotated between a first position (FIG. 21) in which it is operably disengaged from pusher bar assembly 210 and a second position (FIG. 22) in which it is operably engaged with pusher bar assembly 210. Similarly, second actuator knob 214b can be configured to be rotated between first and second positions in which it is operably disengaged and engaged, respectively, with pusher bar assembly 210.

In various embodiments, as a result of the above, the actuator knobs of a stapling instrument can be selectively engaged with a pusher bar assembly such that, in the event that an actuator knob may come into contact with or about tissue surrounding the surgical site when it is advanced, that actuator knob can remain in its retracted position while another actuator knob can be extended to advance the pusher bar assembly distally. In at least one such embodiment, referring to FIG. 22, first actuator knob 214a can be rotated into its second position such that it can be operably engaged with pusher bar assembly 210 while second actuator knob 214b can remain in its retracted position. Thereafter, referring to FIG. 23, first actuator knob 214a can be advanced distally relative to upper handle 226 and lower handle 228 along first side 201 of surgical stapler 210 in order to motivate pusher assembly 210. In at least one embodiment, first actuator knob 214a can be slid within first slot 227 defined between, or within, upper handle 226 and lower handle 228. In various other circumstances, referring to FIG. 28, first actuator knob 214a can remain in its retracted position while second actuator knob 214b can be rotated into its extended position. Similar to the above, second actuator knob 214b can be advanced distally along second side 203 of stapling instrument 210 to advance pusher bar assembly 210 within second slot 229, for example. In at least one embodiment, both actuator knobs 214 can be extended to advance pusher bar assembly 210 distally. In various alternative embodiments, although not illustrated, a stapling instrument can include more than two actuator knobs which can be selectively utilized to motivate a pusher bar and/or knife blade assembly. In effect, as a result of the above, the actuator knobs of a surgical instrument can be engaged with a pusher bar assembly independently of each other.

In various embodiments, further to the above, the actuator knobs of a stapling instrument can be situated in a first position in which they can be held in position and held out of operative engagement with a pusher bar assembly. In at least one embodiment, referring to FIG. 24, stapling instrument 201 can further include guide member 209 which can be configured to guide actuator knobs 214 as they are rotated between their first and second positions. In various embodiments, referring to FIGS. 24–26, guide member 209 can include guide rails 211 which can be slidably received within grooves 213 of actuator knobs 214 such that, when actuator knobs 214 are rotated, guide member 209 can dictate the path along which the actuator knobs 214 are moved. Furthermore, guide rails 211 and grooves 213 can comprise interlocking features which can cooperatively prevent actuator knobs 214 from being unintentionally displaced proximally and/or distally, for example. In at least one such embodiment, guide member 209 can prevent one or more of actuator knobs 214 from being translated along with pusher bar assembly 210 when pusher bar assembly 210 is advanced distally as described above. In various embodiments, a slight friction or interference fit can be present between guide rails 211 and grooves 213 such that the possibility that actuator knobs 214 may be unintentionally rotated into their extended positions can be reduced. Although not illustrated, the actuator knobs can include guide rails extending therethrough which can be slidably received in grooves within the guide member, for example. In any event, referring to FIG. 25, guide member 209 can include one or more retention members 215 which can be configured to retain guide member 209 in position intermediate upper handle 226 and lower handle 228. Furthermore, referring to FIGS. 24 and 25, guide member 209 can include aperture 217 which can be configured to receive retention pin 219 extending therethrough wherein retention pin 219 can be configured to be engaged with upper handle 226 and/or lower handle 228 to retain guide member 209 in position.

In various embodiments, as actuator knobs 214 are rotated between their first and second positions as described above, grooves 213 can be rotated out of engagement with guide rails 211 and actuator knobs 214 can be operatively engaged with pusher bar assembly 210. In at least one embodiment, referring primarily to FIG. 24, pusher bar assembly 210 can include a first clutch feature, such as slots or grooves 205, for example, and actuator knobs 214 can each include a second clutch feature, such as projections 207, for example, wherein the first and second clutch features can be operatively engaged with each other in order to operatively engage one or more of actuator knobs 214 with pusher bar assembly 210. In at least one such embodiment, projections 207 can be closely received within slots 205 such that, when a force is applied to one or more of actuator knobs 214, the
force can be transmitted to pusher bar assembly 210 through projections 207 and the sidewalls of slots 205. In at least one embodiment, similar to the above, a slight friction or interference fit can be present between projections 207 and slots 205 to hold actuators 214 in their extended position. In any event, although not illustrated, the first clutch feature can include projections extending from the pusher bar assembly which can be configured to be received within recesses or slots within the actuator knobs. In addition to or in lieu of the above, referring to FIG. 24, pusher bar assembly 210 can further include second guide rails 221 which can be configured to be slidably received within slots or grooves 223 within actuator knobs 214, wherein rails 221 and grooves 223 can be configured to guide actuator knobs 214 into their second position and/or transmit forces from actuator knobs 214 to pusher bar assembly 210 once they are in their second position. Similar to guide rails 211, guide rails 221 can be configured to create a slight friction or interference fit with grooves 223 to hold actuator knobs 214 in position. Further to the above, in various embodiments, actuator bar 210 can include post 225 about which actuator knobs 214 can be rotated. In at least one embodiment, actuator knobs 214 can include recesses 227 which can be contoured such that the sidewalls of recesses 227 can closely receive and slide around post 225 and, as a result, post 225 can guide actuator knobs 214 as they are rotated between their first and second positions, for example.

[0169] In various embodiments of the present invention, a stapling instrument can include an actuator knob which can be configured to be selectively advanced along a first side of the stapling instrument and a second side of the stapling instrument. In at least one embodiment, referring to FIGS. 29 and 30, stapling instrument 320 can include an upper handle 326, a lower handle 328, and an actuator knob 314, wherein actuator knob 314 can, similar to the above, be configured to advance a pusher bar assembly within a staple cartridge. In at least one embodiment, upper handle 326 and lower handle 328 can define first slot 327 and second slot 329 therebetween, wherein slots 327 and 329 can both be configured to permit actuator knob 314 to slide therethrough. More particularly, in various embodiments, actuator knob 314 can be configured such that it can be selectively slid through first slot 327 along first side 301 or, alternatively, through second slot 329 along second side 303. In various embodiments, referring to FIG. 31, stapling instrument 320 can further include third slot 331 which can be configured to allow actuator knob 314 to be moved from one side of the stapling instrument to the other. In at least one such embodiment, as a result, a surgeon can selectively position actuator knob 314 such that, if it appears that actuator knob 314 may impinge on tissue if it is advanced distally on one side of the stapling instrument, actuator knob 314 can be rotated over to the other side of the stapling instrument before it is advanced. Although the first and second sides of the illustrated embodiment are located on opposite sides of surgical instrument 320, other embodiments are envisioned where the first and second slots, for example, are located on adjacent sides and/or sides which are not directly opposite to each other. Furthermore, other embodiments are envisioned in which the sides of a stapling instrument are not readily discernable, such as instruments having round and/or arcuate portions.

[0170] In various embodiments, referring primarily to FIG. 29, first slot 327 can be configured such that it defines a path for actuator knob 314 which is parallel to, or at least substantially parallel to, a path defined by second slot 329. In at least one embodiment, third slot 331 can be configured to connect first slot 327 and second slot 329 such that it can define a path for actuator knob 314 which is substantially parallel to, or at least substantially perpendicular to, the paths defined by slots 327 and 329. In such embodiments, actuator knob 314 can be rotated over the top of the surgical instrument to move actuator knob 314 from first side 301 to second side 303. In the event that a surgeon decides to reposition actuator knob 314 on first side 301, the surgeon can move actuator knob 314 back through slot 311 until it is positioned within first slot 327 once again. In various alternative embodiments, although not illustrated, a third slot can define a path for actuator knob 314 which is parallel to, or at least substantially parallel to, and/or co-planar with, or at least substantially co-planar with, the paths defined by slots 327 and 329. In further various embodiments, a third slot can define a path which is skew with respect to the paths defined by slots 327 and 329. In any event, a third slot can be configured connect first and second slots such that an actuator knob can be slid therewithin.

[0171] As outlined above, stapling instrument 320 can include a pusher bar assembly which can be operably engaged with actuator knob 314, for example, such that actuator knob 314 can be configured to advance the pusher bar assembly distally. In various embodiments, referring to FIG. 33, stapling instrument 320 can include pusher bar assembly 310 which can include a first portion 333 operably engaged with a handle assembly, for example, and, in addition, a second portion 335 which can be rotatably mounted to first portion 333. In at least one embodiment, first portion 333 can define an axis 337 about which second portion 335 can be rotated. In at least one such embodiment, second portion 335 can include aperture 339 defined therein which can be configured to closely receive first portion 333. In at least one embodiment, although not illustrated, pusher bar assembly 310 can further include one or more retaining members, such as set screws, for example, configured to extend into a groove in first portion 333, for example, for retaining second portion 335 to first portion 333. In various embodiments, second portion 335 can include mount 341 extending therefrom which can be configured to retain actuator knob 314 to second portion 335. In order to move actuator knob from a first side of stapling instrument 320 to the another side, as described above, actuator knob 314 and second portion 335 can be rotated relative to first portion 333 such that actuator knob 314 can be selectively positioned within first slot 327 and second slot 329. In at least one embodiment, although not illustrated, a stapling instrument can have more than two slots for receiving an actuator knob when it is advanced within a staple cartridge. In any event, in various alternative embodiments, first portion 333 and second portion 335 can be fixedly mounted together such that they are rotated together about axis 337. In at least one such embodiment, first portion 333 can be configured to rotate relative to a substantially non-rotatable portion of pusher bar assembly 310.

[0172] Referring to FIG. 34, a surgical stapling instrument, generally 1100, can comprise a first handle portion 1102 and a second handle portion 1104. In various embodiments, first handle portion 1102 and second handle portion 1104 can be configured to be grasped by a surgeon, for example, and can comprise hand grip portion 1106. In at least one embodiment, first handle portion 1102, referring to FIGS. 35 and 36, can include a first cover 1108 attached to a first frame 1110 and, similarly, second handle portion 1104 can include a second
cover 1112 attached to a second frame 1114. Covers 1108 and 1112 can be ergonomically contoured, or otherwise suitably contoured, to assist a surgeon in manipulating stapling instrument 1100 within a surgical site. In various embodiments, handle covers 1108 and 1112, for example, can include enlarged protrusions 1109 and 1113, respectively, which can facilitate the insertion of stapling instrument 1100 into a surgical site. In various embodiments, handle covers 1108 and 1112 can be made of plastic, lightweight materials, and/or any other suitable material, for example, while handle frames 1110 and 1114 can be made of stainless steel, titanium, and/or any other suitable material, for example.

In various embodiments, referring again to FIGS. 34-37, the distal ends of handle portions 1102 and 1104 can comprise an end-effector 1120 which can be configured to treat tissue within a surgical site, for example. In at least one such embodiment, end-effector 1120 can include a staple cartridge channel 1122 configured to receive and/or retain a staple cartridge as described in greater detail further below. In certain embodiments, staple cartridge channel 1122 can comprise a one-piece elongated channel-shaped frame extending from first handle portion frame 1110. In at least one embodiment, staple cartridge channel 1122 can include a pair of opposed, elongated side walls 1124 connected by a bottom wall 1126. Along the rearward, or proximal, portion of staple cartridge channel 1122, a pair of spaced, upstanding side flanges 1128 can extend upwardly from opposed side walls 1124. In various embodiments, the width of staple cartridge channel 1122 between side flanges 1128 can be greater than the width of the upper jaw member, or anvil, 1130 extending from second handle portion 1104. In at least one embodiment, the distance between flanges 1128 can be configured to permit at least a portion of anvil 1130 to be received between side flanges 1128 when the stapling instrument is assembled for operation. As shown in FIG. 35, each side flange 1128 of can include a notch, or recess, 1127, for example, which can be configured to receive one or more latch projections 1131, for example, extending from anvil 1130, and/or any other suitable portion of second handle portion 1104, as described in greater detail further below.

As indicated above, referring once again to FIGS. 34-37, staple cartridge channel 1122 can be configured to support and/or retain a staple cartridge, such as staple cartridge 1150, for example, within end-effector 1120, wherein the staple cartridge can include one or more staples (not illustrated) removably stored therein. In various embodiments, referring to FIGS. 41-43, staple cartridge 1150 can include one or more staple cavities 1151 which can be configured to store staples in any suitable arrangement, such as in at least two laterally-spaced longitudinal rows, for example. In at least one embodiment, referring to FIGS. 42 and 43, staple cartridge 1150 can include staple cartridge body 1152 and pan, or retainer, 1154, wherein staple cartridge body 1152 and/or pan 1154 can be configured to define a channel, or path, for slidably receiving a staple sled and/or cutting member therein. In at least one embodiment, pan 1154 can include flexible arms 1155, for example, which can be configured to engage staple cartridge body 1152 in a snap-fit and/or press-fit arrangement. Referring to FIGS. 43-45, staple cartridge 1150 can further include staple sled assembly 1160 which can include staple sled portion 1162 and, in addition, cutting member 1164. In various embodiments, cutting member 1164 can include cutting edge 1165 and lock arm 1166, for example, wherein lock arm 1166 can be configured to be press-fit and/or snap-fit into aperture 1163 in staple sled 1162 when cutting member 1164 is assembled to staple sled portion 1162. In other various embodiments, staple sled portion 1162 can be integrally molded to cutting member 1164.

Further to the above, referring to FIGS. 41-43, staple cartridge body 1152 can include a slot, such as slot 1156, for example, which can be configured to receive at least a portion of cutting member 1164 therein, and/or any other portion of staple sled assembly 1160 and pusher bar assembly 1200 (discussed below), wherein slot 1156 can be configured to permit cutting member 1164 to be moved between first and second positions within staple cartridge 1150. In various embodiments, slot 1156 can be configured to permit cutting member 1164 to be moved between a proximal position (FIG. 42) and a distal position in order to incise or staple intermediate staple cartridge 1150 and anvil 1130, for example. Referring again to FIGS. 43-45, staple sled portion 1162 can include cam, ramp, or actuator, surfaces 1167 which can be configured to engage staple drivers positioned within staple cartridge 1150. In various embodiments, referring to FIG. 42, staple cartridge 1150 can include staple drivers 1168 which can be lifted, or slid, upwardly within staple cavities 1151 by sled portion 1162 such that the upward movement of staple drivers 1168 can eject, or deploy, staples at least partially positioned within staple cavities 1151. While staple drivers 1168 can be, in fact, lifted vertically upwardly, the term upward, and the like, can mean that staple drivers 1168, for example, are moved toward the top surface, or deck, 1156 of the staple cartridge and/or toward anvil 1130, for example. In certain embodiments, as illustrated in FIG. 42, each staple driver 1168 can include one or more sloped surfaces 1169 oriented at the same angle as a cam surface 1167, and/or any other suitable angle, which can provide a relatively flat, or at least substantially flat, sliding contact surface between staple sled 1162 and staple drivers 1168. In various embodiments, a staple driver can be configured to deploy only one staple, while, in certain embodiments, a staple driver can be configured to simultaneously deploy two or more staples located in adjacent rows, for example. Other devices are disclosed in U.S. patent application Ser. No. 12/030,424, entitled SURGICAL STAPLING INSTRUMENT WITH IMPROVED FIRING TRIGGER ARRANGEMENT, which was filed on Feb. 13, 2008, the entire disclosure of which is incorporated by reference herein.

In various embodiments, as described above, a surgical stapling instrument can include a cutting member/staple sled assembly configured to incise tissue and deploy staples from a staple cartridge. In certain embodiments, though, a surgical stapling instrument may not require, or include, a cutting member. In at least one such embodiment, a staple cartridge can include a staple sled positioned therein and/or a surgical instrument can be configured to move a staple sled into a staple cartridge in order to staple tissue, for example, without otherwise dissecting it. In certain other embodiments, a staple cartridge can include a staple sled positioned wherein a surgical instrument can include a cutting member movable into, or relative to, the staple cartridge. In at least one such embodiment, the cutting member can be advanced into contact with the staple sled such that the cutting member and staple sled can be advanced together. Thereafter, the cutting member can be sufficiently retracted to allow the staple cartridge to be detached from the surgical instrument and replaced with a new staple cartridge having a new staple sled. Such embodiments may be useful when a staple sled
may become worn or deformed during use. Other embodiments are envisioned where a staple cartridge can include a cutting member positioned therein where a surgical instrument can include a staple sled movable into, or relative to, the staple cartridge. In at least one such embodiment, similar to the above, the staple sled can be advanced into contact with the cutting member such that the cutting member and staple sled can be advanced together. Thereafter, the staple sled can be sufficiently retracted to allow the staple cartridge to be detached from the surgical instrument and replaced with a new staple cartridge having a new cutting member. Such embodiments may be useful when a cutting member may become worn or deformed during use. In various embodiments, as described in greater detail below, the staple cartridge can include a protective housing or cover configured to prevent, or at least reduce the possibility of, a surgeon or other clinician from touching the cutting member positioned within the staple cartridge while handling the staple cartridge, for example.

[0177] In various embodiments, further to the above, staple cartridge channel 1122 and/or staple cartridge 1150, for example, can include one or more co-operating projections and/or recesses, for example, which can be configured to removably retain staple cartridge 1150 within staple cartridge channel 1122. Once staple cartridge 1150 has been inserted into staple cartridge channel 1122, in various embodiments, the first handle portion 1102 can be assembled to the second handle portion 1104. In other various embodiments, the staple cartridge may be inserted into the staple cartridge channel after the first and second handle portions have been assembled together. In either event, referring to FIGS. 34-41, first handle portion 1102 and second handle portion 1104 can include proximal ends 1103 and 1105, respectively, which can be assembled together such that the first and second handle portions can be rotatably or pivotably coupled to one another. In various embodiments, referring to FIGS. 35 and 36, first handle portion 1102 can include one or more pins, or projections, 1111 extending therefrom which can be configured to be slidable received within one or more grooves, channels, or slots 1115 in second handle portion 1104. In certain embodiments, slots 1115 can be defined in second handle frame 1114 and projections 1111 can extend from a proximal end post 1107 extending from first handle frame 1110, for example. In order to assemble first handle portion 1102 and second handle portion 1104, referring to FIG. 37, the open ends of slots 1115 can be aligned with projections 1111 such that second handle portion 1104, for example, can be translated relative to first handle portion 1102 and projections 1111 can be slid within slots 1115. In at least one embodiment, as illustrated in FIGS. 35 and 36, the open ends of slots 1115 can be located proximally with respect to their closed ends. In at least one such embodiment, proximal end 1105 of second handle portion 1104 can be positioned distally with respect to proximal end 1103 of first handle portion 1102 such that second handle portion 1104 can be moved proximally in order to position projections 1111 within slots 1115. In various other circumstances, first handle portion 1102 can be positioned proximally with respect to second handle portion 1104 and slid distally in order to position projections 1111 within slots 1115.

[0178] In various embodiments, referring to FIG. 38, second handle portion 1104 can be rotated toward first handle portion 1102 such that anvil 1130 can be moved into position relative to staple cartridge 1150 and/or staple cartridge channel 1122. In certain embodiments, first handle portion 1102 can be rotated toward second handle portion 1104 and/or the first and second handle portions can be rotated toward each other. In any event, projections 1111 and slots 1115, when engaged with one another, can comprise a pivot about which one or both of the first and second handle portions can be moved relative to each other. In various embodiments, second handle portion 1104 can be moved relative to first handle portion 1102 such that anvil 1130 is moved into close opposition to staple cartridge 1150. In certain embodiments, referring to FIG. 39, second handle portion 1104 can be moved relative to first handle portion 1102 such that anvil projections 1131 extending from second handle portion 1104 can be aligned with and/or inserted into recesses 1127 within first handle portion 1102. In various embodiments, referring primarily to FIGS. 35 and 36, first handle portion 1102 can further include latching mechanism 1180 rotatably mounted thereto which can be utilized to engage latch projections 1131 extending from second handle portion 1104 and secure the first and second handle portions together. Although not illustrated, other embodiments are envisioned in which a latching mechanism is rotatably mounted to the second handle portion and latch projections can extend from the first handle portion. In any event, in at least one embodiment, latching mechanism 1180 can be mounted to first frame 1110 by one or more pivot pins 1182 which can be configured to define an axis about which latch 1180 can be rotated.

[0179] In certain embodiments, referring now to FIGS. 37 and 38, latching mechanism 1180 can include latch frame 1184 and, in addition, latch cover 1186 assembled to latch frame 1184. In other various embodiments, the latch cover and the latch frame can comprise an integral unit or, in certain embodiments, the latching mechanism may not even include a cover. In certain embodiments, latch frame 1184 can be channel-shaped and can include a pair of opposed, elongated side walls 1185 which are spaced apart by a distance sufficient to span first frame portion 1110. In at least one embodiment, latch cover 1186 can be made of plastic, lightweight materials, and/or any other suitable materials, for example, while latch frame 1184 can be made of stainless steel and/or any other suitable material, for example. In certain embodiments, when latching mechanism 1180 is closed, as illustrated in FIG. 40, latch cover 1186 can be aligned with first handle cover 1108. Latch cover 1186 can include contoured portion 1187 which can be configured to assist a surgeon in manipulating surgical instrument 1100 wherein, in at least one embodiment, contoured portion 1187 can be aligned with, or at least substantially aligned with, protrusion 1190 extending from first handle cover 1108. Latching mechanism 1180 can further include one or more latch arms 1188 extending therefrom which can be configured to engage one or more latch projections 1131 extending from second handle portion 1104 and pull and/or secure projections 1131 within recesses 1127 as illustrated in FIG. 40. In at least one embodiment, at least one of latch arms 1188 can be integrally-formed with latch frame 1184. In certain embodiments, referring to FIG. 39, at least one of latch arms 1188 can include a distal hook 1189 which can be configured to wrap around at least a portion of projections 1131 so as to encompass or surround, or at least partially encompass or surround, projections 1131. In at least one embodiment, latch arms 1188 can act as an over-center latch to maintain latching mechanism 1180 in its latched, or closed, position.
In use, in various circumstances, one of the first handle portion 1102 and the second handle portion 1104 can be positioned on a first side of tissue within a surgical site and the other handle portion can be rotated into position on the opposite side of the tissue. In such embodiments, staple cartridge 1150 can be positioned on one side of the tissue and anvil 1130 can be positioned on the other side of the tissue. Thereafter, as also outlined above, latching mechanism 1180 can be actuated such that it can be moved between an open position and a closed position in order to latch second handle portion 1104 to first handle portion 1102 and apply a clamping force to the tissue positioned between staple cartridge 1150 and anvil 1130. In certain circumstances, latching mechanism 1180 can be moved between an open position (FIG. 38), a partially-closed, or intermediate, position (FIG. 39), and a closed position (FIG. 40). In at least one such embodiment, referring to FIGS. 38 and 39, latching mechanism 1180 can be moved between an open position in which latch arms 1188 are not engaged with projections 1131 and a partially-closed position in which latch arms 1188 are engaged with projections 1131 such that, although anvil 1130 has been at least partially brought into opposition to staple cartridge 1150, a sufficient gap can remain between anvil 1130 and staple cartridge 1150 which can allow end-effector 1120 to be repositioned relative to the tissue, for example. Once the anvil 1130 and staple cartridge 1150 have been sufficiently positioned relative to the tissue, latching mechanism 1180 can be moved between its partially-closed position and a closed position, as illustrated in FIG. 40.

In various embodiments, further to the above, a surgical stapling instrument can further include a biasing member which can be configured to bias the first handle portion of a stapling instrument away from a second handle portion. In at least one embodiment, as described in greater detail further below, a spring, and/or any suitably resilient material, can be positioned intermediate the first and second handle portions such that the anvil and staple cartridge of the stapling instrument can be biased away from each other. In certain embodiments, the spring can be configured to at least partially separate the first and second handle portions such that a gap exists between the anvil and staple cartridge, wherein the gap can be sufficient to allow tissue to be positioned therebetween. In use, a surgeon can position such a surgical stapling instrument without having to separate and hold the first and second handle portions apart from each other. Such an instrument may be especially useful when the stapling instrument is in a partially-closed configuration and the surgeon is manipulating the instrument within a surgical site. After the surgeon is satisfied with the positioning of the stapling instrument, the surgeon can compress and/or disengage the spring and place the stapling instrument in a closed configuration.

In various circumstances, as outlined above, the distal end of first handle portion 1102 can be moved relative to the distal end of second handle portion 1104, especially when latching mechanism 1180 is not engaged with, or only partially engaged with, projections 1131 of second handle portion 1104. In such circumstances, projections 1111 and slots 1115 at the proximal ends of the first and second handle portions can be configured to retain at least the proximal ends of the first and second handle portions together when the distal ends of the first and second handle portions are being moved relative to each other, for example. Stated another way, projections 1111 and slots 1115 can cooperate to prevent, or at least inhibit, first handle portion 1102 from becoming completely detached from second handle portion 1104. In certain embodiments, a first handle portion can include a first lock portion and a second handle portion can include a second lock portion, wherein the first and second lock portions can be configured to be engaged with one another and prevent the first handle portion from becoming completely detached from the second handle portion. In at least one embodiment, projections 1111 can comprise the first lock portion and slots 1115 can comprise the second lock portion. Previous stapling instruments lacked such lock portions and instead relied on a sole latching mechanism to keep the first and second handle portions together. In circumstances where the latching mechanisms of these previous stapling instruments were not fully engaged with both of the first and second handle portions, the first and second handle portions could become completely detached from one another, thereby requiring a surgeon, for example, to reposition and reassemble the handle portions. In certain circumstances, a complete detachment of the first and second handle portions of these previous staples could expose at least a portion of a cutting member.

In various embodiments, as outlined above, latching mechanism 1180 can be configured to be moved between an open position, a partially-closed position, and a closed position. When latching mechanism 1180 is in its open position, as also outlined above, projections 1111 can be inserted into and/or removed from slots 1115. When latching mechanism 1180 is in its partially-closed position, referring to FIG. 39, latch arms 1188 can be configured to engage latch projections 1131 such that projections 1111 cannot be removed from slots 1115. In at least one such embodiment, latch arms 1188 and latch projections 1131 can be configured to prevent, or at least inhibit, second handle portion 1104 from being moved distally with respect to first handle portion 1102 and, as a result, prevent, or at least inhibit, projections 1111 from being disengaged from slots 1115. Correspondingly, latch arms 1188 and latch projections 1131 can be configured to prevent first handle portion 1102 from being moved proximally with respect to second handle portion 1104. Similar to the above, in various embodiments, latch arms 1188 and latch projections 1131 can also be configured to prevent, or at least inhibit, projections 1111 from being removed from slots 1115 when latching mechanism 1180 is in its closed position (FIG. 40). In certain embodiments, further to the above, latch projections 1131 can extend from second handle portion 1104 at a location which is intermediate its proximal and distal ends. In at least one such embodiment, projections 1111 and slots 1115 can be configured to hold the first and second handle portions together at their proximal ends while latching mechanism 1180 can be utilized to hold the first and second handle portions together at an intermediate location. In any event, in certain embodiments, the first and second handle portions cannot be disengaged from one another unless latching mechanism 1180 is moved into its fully open position. In at least one such embodiment, projections 1111 and slots 1115 cannot be disengaged from one another when latching mechanism 1180 is in a closed and/or partially-closed position.

Once anvil 1130 and staple cartridge 1150 have been sufficiently positioned, the tissue positioned intermediate anvil 1130 and staple cartridge 1150 can be stapled and/or incised. In various embodiments, referring to FIG. 36, surgical stapling instrument 1100 can further include pusher bar assembly 1200 which can be configured to advance and/or retract staple sled assembly 1160 within staple cartridge
1150, for example. In at least one embodiment, pusher bar assembly 1200 can include pusher bar 1202 and firing actuator 1204, wherein firing actuator 1204 can be configured to move pusher bar 1202 and staple sled assembly 1160 distally to deploy staples from staple cartridge 1150 and deform the staples against anvil 1130 as described above. In at least one embodiment, referring to FIGS. 44 and 45, staple sled 1162 can include a groove, channel, or slot 1161 which can be configured to receive, and can be operably connected to, a distal end 1201 (FIG. 36) of pusher bar 1202. In certain embodiments, staple sled assembly 1160 can be operably engaged with pusher bar 1202 when staple cartridge 1150 is inserted into staple cartridge channel 1122. In at least one embodiment, distal end 1201 and slot 1161 can include cooperating features which can allow slot 1161 to be assembled in a transverse direction but prevent, or at least inhibit, distal end 1201 and slot 1161 from being disassembled from one another in a proximal direction and/or distal direction. In other embodiments, pusher bar 1202 can be advanced distally before contacting and engaging staple sled assembly 1160. In at least one such embodiment, the staple sled assembly 1160 can remain stationary until contacted by pusher bar 1202. In any event, as outlined above, actuator 1204 can be operably connected to pusher bar 1202 such that a pulling and/or pulling force can be applied to actuator 1204 and transmitted to pusher bar 1202. In certain embodiments, as described in greater detail below, actuator 1204 can be pivotally connected to a proximal end 1203 of pusher bar 1202 such that actuator 1204 can be selectively rotated between at least first and second positions.

Further to the above, referring to FIGS. 34, 46, and 47, actuator 1204 can be movable between a first position on a first side 1116 of surgical stapling instrument 1100 (FIG. 46), a second position on a second side 1117 (FIG. 47), and an intermediate position (FIG. 34) located at the proximal ends 1103 and 1105 of the first and second handle portions 1102 and 1104. Once actuator 1204 has been rotated into position on one of the first and second sides 1116, 1117, actuator 1204 can be advanced distally. In various circumstances, as a result, a surgeon may select whether to move actuator 1204 distally along first side 1116 or second side 1117. Such circumstances may arise when it is more likely that actuator 1204 may impinge on tissue surrounding the surgical site, for example, when actuator 1204 is moved distally along one side of the surgical instrument as compared to the other. In various embodiments, referring to FIGS. 35 and 36, actuator 1204 can include arm 1206 extending therefrom where arm 1206 can be integrally mounted to proximal end 1202 of pusher bar 1202. In certain embodiments, referring once again to FIGS. 34, 46, and 47, surgical instrument 1100 can include a first slot (not illustrated) extending along first side 1116 and a second slot 1118 extending along second side 1117, wherein the first and second slots can be configured to slidably receive at least a portion of actuator 1204. In at least one embodiment, the sidewalls of the first and second slots can confine, or at least assist in confining, the movement of actuator 1204 such that it can be moved along a predetermined path. Referring to FIG. 47, second slot 1118, for example, can be defined between first handle portion 1102 and handle portion 1104 such that, when actuator 1204 is moved distally along second side 1117, arm 1206 of actuator 1204 can slid intermediate the first and second handle portions. Similar to the above, the first slot can also be defined intermediate the first and second handle portions.

Referring again to FIGS. 46 and 47, surgical instrument 1100 can further include intermediate slot 1119 which can also be configured to allow arm 1206, and/or any other suitable portion of actuator 1204, to slide therein. In at least one such embodiment, intermediate slot 1119 can connect the first and second slots such that, when actuator 1204 is positioned in its intermediate position, actuator 1204 can be moved into either one of its first and second positions. In certain embodiments, the first slot, second slot 1117, and intermediate slot 1119 can be parallel, or at least substantially parallel, to one another and/or lie in the same plane, although other embodiments are envisioned in which one or more of the slots is not parallel to the others and/or lies in a different plane. Furthermore, although the first and second sides of the illustrated embodiment are located on opposite sides of surgical instrument 1100, other embodiments are envisioned where the first and second slots, for example, are located on adjacent sides and/or sides which are not directly opposite to each other. Furthermore, other embodiments are envisioned in which the sides of a stapling instrument are not readily discernable, such as instruments having round and/or arcuate portions.

In various embodiments, further to the above, surgical stapling instrument 1100 can further include a locking mechanism which can prevent, or at least inhibit, actuator 1204 and, correspondingly, staple sled assembly 1160, from being advanced prematurely. In at least one embodiment, the locking mechanism can be configured to prevent, or at least inhibit, actuator 1204 from being advanced distally prior to latching mechanism 1180 being moved into a closed, or an at least partially-closed, position. In certain embodiments, generally referring to FIG. 38, surgical stapling instrument 1100 can further including locking mechanism 1220 which can be engaged with actuator 1204 and can remain engaged with actuator 1204 while latching mechanism 1180 is in a fully open position (FIG. 38) and/or at least substantially-open position. In various embodiments, locking mechanism 1220 can include lock 1222 which can be biased into engagement with actuator 1204 by a biasing force applied thereto by lock spring 1224, for example. In at least one such embodiment, actuator 1204 can include one or more grooves, channels, or slots (not illustrated) which can be configured to receive at least a portion of lock 1222. In use, locking mechanism 1220 can hold actuator 1204 in position until latching mechanism 1180 is moved into its fully closed position (FIG. 40) and/or at least substantially-closed position. In such circumstances, in at least one embodiment, locking mechanism 1180 can be configured to engage locking mechanism 1220 and disengage lock 1222 from actuator 1204. In at least one such embodiment, referring to FIGS. 38-40, latching mechanism 1180 can further include cam 1183 which can be configured to engage cam surface 1223 on lock 1222 when latching mechanism 1180 is moved into its closed position and, as a result, slide, and/or otherwise move, lock 1222 away from actuator 1204. In various embodiments, cam 1183 can comprise a wall, rib, and/or ridge extending from latch cover 1186 and/or latch frame 1184. In any event, once lock 1222 has been sufficiently disengaged from actuator 1204, in at least one embodiment, actuator 1204 can be moved from its intermediate position, illustrated in FIG. 34, into one of its first and second positions, as illustrated in FIGS. 46 and 47.

As described above, locking mechanism 1220 can be configured to prevent, or at least inhibit, drive bar 1202 from being advanced distally prior to latching mechanism 1180 being moved into a predetermined position, such as, for
example, a closed position and/or partially-closed position. Advantageously, locking mechanism 1220 may also prevent, or at least inhibit, staple sled assembly 1160 from being advanced prior to the first handle portion 1102 and the second handle portion 1104 being assembled together. In effect, locking mechanism 1220 can prevent tissue positioned intermediate unvil 1130 and staple cartridge 1150 from being cut and/or stapled prior to unvil 1130 and staple cartridge 1150 being properly positioned relative to the tissue. Also, in effect, locking mechanism 1220 can prevent staples from being deployed into the tissue prior to an appropriate clamping force being applied to the tissue. In any event, when latching mechanism 1180 is returned to its fully open position, and/or a partially-open position, cam 1183 can be moved away from lock 1222 such that lock spring 1124 can be engaged with actuator 1204 once again. In various other embodiments, referring to FIGS. 38 and 39, locking mechanism 1220 can include a lock 1222 comprising a cam surface 1223 and, in addition, a step 1226 which can limit the relative movement of lock 1222. In at least one embodiment, cam 1183, for example, can be configured to contact cam surface 1223 and, owing to the contoured, beveled, and/or angled surface of cam surface 1223, cam 1183 can be configured to drive lock 1222 distally as illustrated in FIG. 49. Lock 1222 can be driven distally such that pin 1228, which extends from lock 1222, can be moved between a first position (FIG. 48) in which it is positioned within aperture 1229 in actuator 1204 and a second position (FIG. 49) in which pin 1228 has been sufficiently removed from aperture 1229. In various embodiments, stop 1226 can be configured such that, as lock 1222 is driven distally, step 1226 can come into contact with cam 1183 once lock 1222 has been sufficiently displaced. In such embodiments, stop 1226 can be configured to control the second, or displaced, position of lock 1222. Similar to the above, as actuator 1180 is moved out of its closed position and cam 1183 is disengaged from locking mechanism 1220, lock spring 1224 can move lock 1222 into engagement with actuator 1204 once again.

[0189] In various embodiments, as described above, cutting member 1165 can be at least partially positioned within slot, or channel, 1156 and, as illustrated in FIG. 43, at least the upper, or top, portion of cutting member 1164 can extend above deck 1158. In at least one embodiment, referring to FIGS. 41-43, housing 1170 can include a first wall, or portion, 1172 extending from a first portion 1157 of staple cartridge body 1152, a second wall, or portion, 1174 extending from a second portion 1159 of staple cartridge body 1152, and a top wall, or portion, 1176 extending between first wall 1172 and second wall 1174. In certain embodiments, a housing may comprise only one support wall, or support portion, extending from a staple cartridge body and, in addition, a top wall, or top portion, extending therefrom. In other embodiments, a housing may comprise one or more substantially planar, interior surfaces and/or top wall. In at least one such embodiment, the side walls of the housing can be configured such that they extend above the top of the cutting member, or at least extend above a cutting edge of the cutting member, for example. In any event, as illustrated in FIG. 43, at least a portion of cutting member 1164 can be positioned underneath top wall 1176 and/or between side walls 1172 and 1174 when staple sled assembly 1160 is in its proximal position. In certain embodiments, cutting member 1164 can be entirely positioned underneath top wall 1176, and/or entirely positioned within housing 1170. In at least one embodiment, cutting member 1164 can be positioned underneath top wall 1176 such that staple surface 1165 does not extend beyond the distal edge 1175 and/or the proximal edge 1177 of top wall 1176. In at least one embodiment, housing 1170 can include a rear wall 1178 which can be configured to limit the proximal movement of cutting member 1164 and/or any other portion of staple sled assembly 1160. In various embodiments, at least a portion of housing 1170, for example, can be integrally-formed with staple cartridge body 1152. In at least one such embodiment, first wall 1172, second wall 1174, top wall 1176, and/or rear wall 1178 can be formed when staple cartridge body 1152 is injection molded, for example. In certain embodiments, at least a portion of housing 1170 can be assembled to staple cartridge body 1152 via a snap-fit arrangement, press-fit arrangement, and/or any other suitable manner.

[0190] In various embodiments, further to the above, cutting member 1164 can be defined by a planar, or an at least substantially planar, body having a knife edge extending along at least one side of the cutting member body. In at least one such embodiment, first wall 1172 and/or second wall 1174 can be configured and arranged such that they can include planar, or at least substantially planar, interior surfaces 1173 which are parallel, or at least substantially parallel, to the side surfaces of cutting member 1164. In certain embodiments, cutting member 1164 can be closely received between the interior surfaces 1173 of walls 1172 and 1174. In at least one such embodiment, the distance between walls 1172 and 1174 may be the same as, or at least substantially the same as, the width of slot 1156. In any event, a housing can be configured such that at least a portion of the housing extends over at least a portion of slot 1156, for example. In certain embodiments, housing 1170 can completely enclose or surround a cutting member 1164 and/or cutting surface 1165. In at least one embodiment, although not illustrated, a housing can include a break-away and/or incises portion which can be at least partially detached, separated, and/or otherwise deformed in order to permit a cutting member to exit the housing. In at least one such embodiment, the tissue cutting
surface can be configured to contact the housing to break and/or incise a housing wall, for example. In various embodiments, the housing wall can include a thin portion, a reduced-thickness portion, score mark, and/or any other configuration to facilitate the deformation and/or incision of the housing wall. In certain embodiments, a cutting member can include one or more additional cutting surfaces and/or anvils, for example, which can be configured to deform and/or incise the housing. In at least one embodiment, the housing can include a movable and/or flexible portion, such as a hinged member and/or flexible flap, for example, which can be configured to sufficiently move and/or flex to allow the cutting member to pass thereby. In any event, embodiments are envisioned in which the cutting member can have any suitable configuration for incising tissue and the protective housing can have any suitable configuration for at least partially enclosing or surrounding the cutting member. Furthermore, although a cutting member can comprise a sharpened edge as described above, other suitable cutting members are envisioned, such as those supplied with an electrical current sufficient to dissect tissue, for example.

[0191] As described above, housing 1170 can be configured to at least partially cover, enclose, and/or surround a cutting member when it is in its proximal position. In various embodiments, the cutting member can be advanced distally to incise tissue, for example, and then retracted proximally in order to position the cutting member within housing 1170 once again. In such embodiments, the cutting member can be at least partially covered by housing 1170 when the staple cartridge is assembled to and removed from a surgical stapling instrument. In certain embodiments, a new, or unspent, staple cartridge can be inserted into the staple cartridge channel to replace the at least partially spent staple cartridge. In at least one such embodiment, the new staple cartridge can include a new cutting member and/or staple sled assembly positioned therein, although embodiments are envisioned in which the previously-used cutting member and/or staple sled assembly can be sufficiently withdrawn from the spent staple cartridge and advanced into the new staple cartridge in order to be re-used once again. In embodiments where a new cutting member and/or staple sled assembly is provided with each new staple cartridge, a sharp cutting edge, for example, can be utilized with each staple cartridge.

[0192] In various embodiments, although not illustrated, a staple cartridge can include two or more housings configured to at least partially cover a cutting member when it is in two or more positions. In at least one embodiment, a staple cartridge can include a proximal housing configured to at least partially cover the cutting member when it is in a proximal position, for example, and, in addition, a distal housing configured to at least partially cover the cutting member when it is in a distal position, for example. In at least one such embodiment, the cutting member can be positioned within the proximal housing when the staple cartridge is assembled to a surgical stapling instrument and, in certain embodiments, the cutting member can be advanced into the distal housing after it has transected tissue positioned within the end-effector, for example. In such embodiments, as a result, the cutting member can be at least partially positioned within the distal housing when the staple cartridge is removed from the surgical stapler. Such embodiments may be particularly useful when a vessel, for example, is positioned intermediate the proximal housing and the distal housing of the staple cartridge. In various embodiments, although not illustrated, a cutting member can be moved proximally from a distal position to a proximal position, and/or any other suitable position.

[0193] In various embodiments, as discussed above, staple cartridge 1150 can be inserted into staple cartridge channel 1122. Referring now to FIG. 92, a proximal end 1213 of staple cartridge 1150 can be positioned within a proximal end 1123 of staple cartridge channel 1122 while a distal end 1211 of staple cartridge 1150 can be positioned within a distal end 1121 of staple cartridge channel 1122. In at least one embodiment, the distal end 1121 of staple cartridge channel 1122 can comprise one or more projections and/or one or more recesses which can be correspondingly aligned with one or more projections and/or one or more recesses in the distal end 1211 of staple cartridge 1150, for example. In at least one such embodiment, staple cartridge channel 1122 can comprise a projection, or tab, 1279 and a recess, or slot, 1278, wherein each side of staple cartridge 1150 can comprise, referring to FIG. 95, a projection 1274 configured to be positioned within a recess 1278 and, in addition, a recess 1270 configured to receive a projection 1279. In various embodiments, each recess 1270 of staple cartridge 1150 can comprise opposing sidewalls 1272 and 1273 and a distal surface 1271, wherein the distal surface 1271 can be positioned against the projection 1279 positioned therein when the staple cartridge 1150 is positioned in staple cartridge channel 1122. In various circumstances, as discussed in greater detail below, the distal surfaces 1271 of recesses 1270 can serve as a datum surface from which certain features of the staple cartridge 1150 can be predetermined. In some circumstances, the distal end 1211 of staple cartridge 1150 can be aligned with and/or inserted into the distal end 1121 of staple cartridge channel 1122 before the proximal end 1213 of staple cartridge 1150 is inserted into the proximal end 1123 of staple cartridge channel 1122. For example, the distal end 1211 of staple cartridge channel 1150 can be aligned with the staple cartridge channel 1122 such that projections 1279 are positioned within recesses 1270 wherein, thereafter, the staple cartridge 1150 can be rocked, or rotated, toward staple cartridge channel 1122 such that proximal end 1213 of staple cartridge 1150 is inserted into the proximal end 1123 of staple cartridge channel 1122.

[0194] When distal end 1211 of staple cartridge 1150 is engaged with the distal end 1121 of staple cartridge channel 1122, as described above, the projections 1274 of staple cartridge 1150 can be inserted into the recesses 1279 of staple cartridge channel 1122 by hooking the projections 1274 underneath the projections 1278 of staple cartridge channel 1122. In such circumstances, the co-ordination of projections 1274 and 1278 and recesses 1270 and 1279 can attach the distal end 1211 of staple cartridge 1150 to the distal end of staple cartridge 1122 and, in addition, align the staple cartridge 1150 with the staple cartridge channel 1122 such that the staple cartridge 1150 can be inserted between the sidewalls 1124 of staple cartridge channel 1122. Once the distal end 1211 of staple cartridge 1150 has been hooked to staple cartridge channel 1122, at least one of the staple cartridge 1150 and the staple cartridge channel 1122 can be rotated toward the other. In various circumstances, referring again to FIGS. 92 and 95, the staple cartridge 1150 can be pivoted toward the staple cartridge channel 1122 such that alignment slots 1280 in staple cartridge channel 1150 become aligned with side flanges 1128. In various embodiments, the staple cartridge 1150 can comprise alignment slots 1280 on opposite sides thereof which can each be configured to receive a
side flange 1128. In at least one embodiment, each alignment slot 1280 can comprise lateral sidewalls 1283 and 1284 and a basewall 1281 extending between the sidewalls 1283 and 1284. Further to the above, a predetermined distance 1289 can be measured between the distal datum surfaces 1271 of recesses 1270 to the distal basewalls 1281 of alignment slots 1280. Referring now to FIGS. 93 and 95, the predetermined distance 1288 between the distal end of the projections 1279 and the distal end of the side flanges 1128 can be such that it is shorter than the distance 1289 between the distal surfaces 1271 of recesses 1270 and the basewalls 1281 of alignment slots 1280. Owing to the distance 1288 being shorter than the distance 1289, the staple cartridge 1150 can be rotated into position as described above such that side flanges 1128 can enter into alignment slots 1280. In various embodiments, alignment slots 1280 can be sized and configured such that the side flanges 1128 are closely received between the sidewalls 1283 and 1284 such that there is little, if any, relative movement between the side flanges 1128 and the sidewalls of the alignment slots 1280, for example.

In various alternative embodiments, further to the above, the proximal end 1213 of staple cartridge 1150 can be inserted into the distal end 1121 of staple cartridge channel 1122 and slid proximally between sidewalls 1124 such that the proximal end 1213 of staple cartridge channel 1150 enters into the proximal end 1123 of staple cartridge channel 1122. During such sliding movement, the side flanges 1128 can enter into alignment slots 1280 and, in addition, the projections 1279 can enter into the recesses 1270. In certain embodiments, the staple cartridge 1150 can be both slid and rotated into the staple cartridge channel 1122. In any event, in various embodiments, the staple cartridge 1150 and the staple cartridge channel 1122 can be configured such that the staple cartridge 1150 can be removably secured within the staple cartridge channel 1122. In at least one embodiment, referring primarily now to FIGS. 95 and 100, the staple cartridge 1150 can comprise one or more retention features which can be configured to releasably engage one or more retention features in the staple cartridge channel 1122. More particularly, in at least one such embodiment, the staple cartridge 1150 can comprise one or more retention slots 1190 which can be configured to engage one or more retention keys 1195 in the staple cartridge channel 1122. In various embodiments, referring again to FIG. 95, each retention slot 1190 can comprise a first, or entrance, portion 1191 which can be configured to receive a retention key 1195 therein and, in addition, a second portion 1192 which can be configured to receive the retention key 1195 after it has passed through the entrance portion 1191. The entrance portion 1191, in certain embodiments, can define a first width between a proximal side 1193 and a distal side 1194 of retention slot 1190 and, in addition, the second portion 1192 can define a second width between the proximal side 1193 and the distal side 1194 which is wider than the first width of entrance portion 1191. In various embodiments, the first width of entrance portion 1191 can be narrower than the width of the retention key 1195 and the second width of second portion 1192 can be wider than the width of the retention key 1195. In at least one such embodiment, a retention slot 1190 can be configured to engage a retention key 1195 in at least one of a pre-fit and/or a snap-fit manner. In certain embodiments, at least one of the proximal side 1193 and/or the distal side 1194 can be configured to flex or splay outwardly as the retention key 1195 is inserted into retention slot 1190. In at least one such embodiment, the proximal sides 1193 can be displaced proximally. In any event, referring to FIG. 100, once the retention slot 1190 has received the retention key 1195, the proximal side 1193 of retention slot 1190 can be positioned on a proximal side 1196 of retention key 1195 and the distal side 1194 of retention slot 1190 can be positioned on a distal side 1197 of retention key 1195.

As outlined above, the staple cartridge 1150 can be assembled into the staple cartridge channel 1122 by coupling the distal end 1211 of staple cartridge 1150 to the distal end 1211 of staple cartridge channel 1122 and then rotating the proximal end 1213 of staple cartridge 1150 into the proximal end 1123 of staple cartridge channel 1122. In at least one such embodiment, the retention slots 1190 can be configured to engage the retention keys 1195 as the staple cartridge 1150 is rotated into its seated position within staple cartridge channel 1122. Referring now to FIG. 93, a predetermined distance 1199 between the distal datum surfaces 1271 of recesses 1270 and the retention slots 1190 can be sized and configured such that the retention slots 1190 are aligned with the retention keys 1195 as the staple cartridge 1150 is rotated into position as described above. Correspondingly, in at least one embodiment, a distance between the distal ends of projections 1279 and retention keys 1195 can be such that it equals, or at least substantially equals, the distance 1199. In various circumstances, the above-mentioned distances can be measured to the center of the features comprising retention slots 1190 and retention keys 1195. For example, the distance 1199 can be measured to a position in the center of slot 1190 intermediate the proximal and distal sidewalls thereof, for example. In various embodiments, the retention slot 1190 can further comprise lead-in, beveled, and/or radius-ed surfaces, which can be configured to guide, or direct, the retention keys 1195 into the retention slots 1190. In at least one such embodiment, these lead-in surfaces can be wider than the first portions 1191.

As staple cartridge 1150 is rotated into staple cartridge 1122, a cutting member and/or staple deploying sled positioned within the staple cartridge 1150 can be operably engaged with the pusher bar 1202. More particularly, referring now to FIGS. 97-99, the staple cartridge 1150 can include a cutting member 1160 which can be operably coupled with pusher bar 1202 such that, after the staple cartridge 1150 has been seated within the staple cartridge channel 1122, the pusher bar 1202 and cutting member 1160 can be advanced together as described above. In at least one embodiment, the cutting member 1160 can comprise a slot 1161 which can be configured to receive a distal drive projection 1294 (FIG. 93) at the distal end of pusher bar 1202. More particularly, referring now to FIG. 101, the slot 1161 of cutting member 1160 can be aligned with an access slot 1290 in the bottom of the staple cartridge 1150 such that, as the proximal end 1213 of staple cartridge 1150 is seated in the proximal end 1123 of staple cartridge channel 1122, the drive projection 1294 of pusher bar 1200 can extend through the access slot 1290 into the slot 1161 of cutting member 1160. In various embodiments, the slot 1161 and the drive projection 1294 can be sized and configured such that there is little, if any, relative movement therebetween. More particularly, referring again to FIGS. 98 and 99, the slot 1161 can comprise a distal sidewall 1291 and a proximal sidewall 1292 wherein the drive projection 1294 can be securely received between the sidewalls 1291 and 1292. In various embodiments, referring again to FIGS. 93 and 101, the pusher bar 1202 can
further comprise a recess, or slot, 1295 positioned proximally with respect to the drive projection 1294 wherein the slot 1295 can be configured to receive a proximal projection 1293 (FIG. 97) extending from the cutting member 1160. Similar to the above, the slot 1295 can be defined by sidewalls which can be configured to closely receive the proximal projection 1293 such that there is little, if any, relative movement therebetween.

[0198] As described above, the slot 1161 of cutting member 1160 can be positioned within the staple cartridge 1150 such that it is aligned with the drive projection 1294 of the push bar 1202 when the staple cartridge 1150 is seated within the staple cartridge channel 1122. Referring now to FIG. 96, a predetermined distance 1299 can be defined between the distal surfaces 1271 of recesses 1270 and the slot 1161, wherein the distance 1299 can be equal to, or at least substantially equal to, a predetermined distance 1297 between the distal end of the projections 1279 and the drive projection 1294. In various circumstances, the cutting member 1160 can be moved through a range of positions between a proximal-most position, in which it is positioned in housing 1170, and a distal-most position after it has been advanced through the cutting slot 1156. In various embodiments, the distance 1299 can be measured with respect to the cutting member 1160 when it is in its proximal-most position. Similar to the above, the distances 1297 and 1299 can be measured to the center or midpoint of the drive projection 1297 and slot 1161, respectively. In various embodiments, the surgical instrument 1100 can further comprise a locking mechanism which can be configured to hold the pusher bar 1202 in position while the cutting member 1160 is engaged with the drive projection 1294. Similar to the above, in certain embodiments, a distance 1298 can be defined between the distal end of projections 1279 and the recess 1295 of the push bar 1202 wherein the distance 1298 can be equal to, or at least substantially equal to, the distance between the distal surface 1271 of recesses 1270 and the projection 1293 of the cutting member 1160. In various embodiments, referring primarily now to FIGS. 97 and 100, the staple cartridge 1150 can comprise a clearance region defined between the proximal end 1295 of the staple cartridge body 1152 and the proximal end 1294 of the staple cartridge pin 1154, wherein such a clearance region can be configured to receive the pusher bar 1202 and/or a portion of the staple cartridge channel 1122 therein, for example. In any event, the pusher bar 1202 can be advanced distally once it has been engaged with the cutting member 1160, wherein such movement is depicted in FIG. 100 which illustrates the distal end 1201 of pusher bar 1202 in a proximal position (illustrated with solid lines) and a second, distal position (illustrated with phantom lines), for example.

[0199] In various embodiments, as described above, the distal end 1211 of staple cartridge 1150 can be engaged with the distal end 1121 of the staple cartridge channel 1122 and then pivoted into staple cartridge channel 1122 such that the proximal end 1213 of staple cartridge 1150 can be seated in the proximal end 1123 of staple cartridge channel 1122. Such a process can comprise engaging the projections 1274 of staple cartridge 1150 under the projection 1276 of staple cartridge channel 1122 and then, as described above, rotating the staple cartridge 1150 until alignment slots 1280 are positioned adjacent to flanges 1182. At such point, in various embodiments, the cutting member 1160 may not be engaged with the pusher bar 1202 and, in addition, the retention slots 1190 may not be engaged with the retention keys 1195. As a result, the surgeon, or clinician, can adjust the position of the staple cartridge 1150 within the staple cartridge channel 1122 before the staple cartridge 1150 is locked into position. Once the side flanges 1182 have been at least partially positioned in alignment slots 1280, the proximal end 1213 can be further rotated toward the staple cartridge channel 1122. At such point, the cutting member 1160 can come into operable engagement with the pusher bar 1202 and, in addition, the retention slots 1190 can engage the retention keys 1195. In various embodiments, the cutting member 1160 can operably engage the pusher bar 1202 at the same time, or at least substantially the same time, as the retention slots engage retention keys 1195. More particularly, in at least one embodiment, the drive projection 1294 of the pusher bar 1202 can enter slot 1161 of cutting member 1160 at the same time that the retention keys 1195 enter into, or snap into, the second portions 1192 of slots 1190. In at least one such embodiment, the cutting member 1160 may not be advanceable by the pusher bar 1202 until the staple cartridge 1150 is snapped into, or seated in, place. In certain alternative embodiments, the cutting member 1160 can be operably engaged with the pusher bar 1202 before the retention keys 1195 are fully seated within the retention slots 1190 when the proximal end 1213 of the staple cartridge 1150 is seated in the proximal end 1123 of the staple cartridge channel 1122. In various embodiments, the retention slots 1190 can be aligned with each other such that they engage the retention keys 1195 at the same time, or at least substantially the same time. In at least one such embodiment, the retention slots can be configured such that the retention keys 1195 enter into the second portions 1192 of the retention slots 1190 simultaneously. In at least one embodiment, the retention slots 1190 can be positioned along an axis which is transverse to or perpendicular to a longitudinal axis defined by the cutting slot 1156. In various embodiments, the retention slots 1190, and the axis defined therebetween, can be positioned proximally with respect to the cutting member 1160 regardless of the position of the cutting member 1160 including when the cutting member 1160 is in its proximal-most position, for example.

[0200] In various embodiments, the cutting slot 1156 can define a first body portion 1152a on a first side thereof and a second body portion 1152b on a second, or opposite, side thereof. Referring to FIGS. 95 and 98, the first body portion 1152a can comprise a first plurality of staple cavities 1151 and, in addition, the second body portion 1152b can comprise a second plurality of staple cavities 1151. In at least one embodiment, the first body portion 1152a can comprise a proximal-most staple cavity 1151a which can be positioned proximally relative to the other staple cavities 1151 in the first body portion 1152a. In at least one embodiment, the entirety of staple cavity 1151a can be positioned proximally relative to base wall 1281 of the alignment slot 1280 in first body portion 1152a, while, in other embodiments, at least a portion of staple cavity 1151a can be positioned proximally relative to the base wall 1281. As illustrated in FIG. 97, the alignment slot 1280 in the first body portion 1152a is positioned laterally with respect to the proximal-most staple cavity 1151a and, in addition, laterally with respect to the cutting slot 1156. Further to the above, the first body portion 1152a can comprise a second proximal-most staple cavity 1151c which can be positioned proximally relative to the other staple cavities 1151 in first body portion 1152a except for proximal-most staple cavity 1151a. In at least one embodiment, the entirety of staple cavity 1151c can be positioned proximally relative to...
base wall 1281 of the alignment slot 1280 in first body portion 1152a, while, in other embodiments, at least a portion of staple cavity 1151c can be positioned proximally relative to the base wall 1281. As illustrated in FIG. 97, the alignment slot 1280 is the first body portion 1152a is at least partially positioned laterally with respect to the second proximal-most staple cavity 1151c. Still referring to FIG. 97, the first body portion 1152a can comprise a retention slot 1190 therein which can be positioned proximally with respect to the staple cavities 1151 therein, including the staple cavities 151a and 1151c, for example.

[0201] Referring to FIG. 95, further to the above, the second body portion 1152b can comprise a proximal-most staple cavity 1151b which can be positioned proximally relative to the other staple cavities 1151 in second body portion 1152. In at least one embodiment, the entirety of staple cavity 1151b can be positioned proximally relative to base wall 1281 of the alignment slot 1280 in second body portion 1152b, while, in other embodiments, at least a portion of staple cavity 1151b can be positioned proximally relative to the base wall 1281. As illustrated in FIG. 95, the alignment slot 1280 in the second body portion 1152b is positioned laterally with respect to the proximal-most staple cavity 1151b and the cutting slot 1156. Further to the above, the second body portion 1152b can comprise a second proximal-most staple cavity 1151d which can be positioned proximally relative to the other staple cavities 1151 in second body portion 1152b except for proximal-most staple cavity 1151b. In at least one embodiment, the entirety of staple cavity 1151d can be positioned proximally relative to base wall 1281 of the alignment slot 1280 in second body portion 1152b, while, in other embodiments, at least a portion of staple cavity 1151d can be positioned proximally relative to the base wall 1281. As illustrated in FIG. 95, the alignment slot 1280 in the second body portion 1152b is at least partially positioned laterally with respect to the second proximal-most staple cavity 1151d. Still referring to FIG. 95, the second body portion 1152b can comprise a retention slot 1190 therein which can be positioned proximally with respect to the staple cavities 1151 therein, including the staple cavities 1151b and 1151d, for example.

[0202] In various embodiments, further to the above, the staple cartridge body 1152 can be comprised of plastic and can be formed utilizing an injection molding process. Thereafter, in various embodiments, the staple drivers 1168 (FIG. 42) can be assembled into staple cavities 1151 and the cutting member 1160 can be positioned within the cartridge body 1152 such that the cutting member 1164 is located within the housing 1170, as described above. The staple cartridge pan 1154 can then be assembled to the staple cartridge body 1152. In various embodiments, referring now to FIG. 96, the distal end 1277 of staple cartridge pan 1154 can be aligned with the proximal end 1295 of the staple cartridge body 1152 such that the staple cartridge body can be slid within the staple cartridge pan 1154 between opposing walls 1154a and 1154b, for example. The staple cartridge body 1152 and pan 1154 can be slid relative to one another until pan projections 1276 are positioned within recesses 1270 and projections 1274 are positioned within pan recesses 1275. At the same time, the lock projections 1288 extending from staple cartridge body 1152 can be received within the lock apertures 1287 in staple cartridge pan 1154 such that pan 1154 can be locked to staple cartridge body 1152. In various embodiments, the sidewalks 1154a and 1154b of pan 1154 can flex or splay outwardly as they pass over lock projections 1288 and then elastically return inwardly when lock apertures 1287 are aligned with lock projections 1288. At such point, the arms 1155 extending from pan 1154 can be aligned with and positioned within the retention slots 1287 in staple cartridge body 1152. In certain embodiments, referring now to FIG. 101, the staple cartridge 1150 can further comprise a retention member, such as retention member 1300, for example, which can be configured to selectively obstruct slot 1301 in staple cartridge body 1152, for example. In at least one embodiment, the retention member 1300 can comprise a pivotable arm 1303 which can be rotated between a first position in which it extends across slot 1301 (illustrated in solid lines) and a second position in which it is positioned adjacent to slot 1301 (illustrated in phantom lines). In at least one such embodiment, an integral pivot pin 1302 (FIG. 95) can extend from arm 1303 into an aperture in staple cartridge body 1152 which can define an axis about which the arm 1303 can be rotated. In certain embodiments, the arm 1303 can include a lock member 1304 extending therefrom which can be configured to be releasably engaged with a lock cavity 1305 in staple cartridge body 1152 in order to hold the arm 1303 in at least one of its first and second positions, for example. In certain embodiments, the positioning of arm 1303 across slot 1301 can prevent, or at least inhibit, the cutting member 1160, for example, from sliding out of the staple cartridge 1150.

[0203] In order to facilitate the insertion and removal of the staple cartridge 1150 from staple cartridge channel 1122, in various embodiments, the staple cartridge 1150 can comprise gripping portions positioned on opposite sides thereof, for example. In at least one embodiment, referring now to FIGS. 97 and 101, the staple cartridge body 1152 can comprise lateral portions 1285 positioned adjacent to alignment slots 1280 wherein the lateral portions 1285 can be gripped and/or pushed on by a clinician in order to seat the proximal end 1213 of staple cartridge 1150 in the proximal end of staple cartridge channel 1122, for example. Such a force can be applied to top, or tissue-contacting, surfaces of the lateral portions 1285 as the proximal end 1213 of staple cartridge 1150 is rotated into position as described above. In various embodiments, a lifting force can be applied to lateral portions 1285 in order to lift the proximal end 1213 of staple cartridge 1150 out of the staple cartridge channel 1122. In at least one such embodiment, referring primarily to FIG. 101, each lateral portion 1285 can comprise one or more steps, ridges, and/or elevations, such as elevations 1287a, 1287b, and/or 1287c, for example, which can be configured to improve the clinician’s grip on the lateral portions 1285. In various embodiments, the elevations 1287a, 1287b, and/or 1287c can be positioned at different heights relative to one another. In any event, the staple cartridge 1150 can be removed from channel 1122 by lifting the proximal end 1213 of staple cartridge 1150 out of channel 1122 and then unhooking, or disengaging, the distal end 1211 of staple cartridge 1150 from the distal end 1211 of channel 1122. For example. As staple cartridge 1150 is removed from the channel 1122, the slot 1161 within cutting member 1160 can be moved away and disengaged from the drive projection 1294 of pusher bar 1202, for example.

[0204] In various circumstances, further to the above, the pusher bar 1202 and cutting member 1160 can be returned to their proximal positions before the staple cartridge 1150 is removed from the staple cartridge channel 1122. In such a position, as described above, the cutting edge 1165 can be
positioned within the housing 1170. In various embodiments, referring now to FIG. 102, an alternative embodiment of a staple cartridge 1150 is depicted without a housing 1170. In at least one such embodiment, the cutting edge 1165 can at least partially extend above the deck surface 1158 of the staple cartridge body 1152 in its proximal position and/or any other distally advanced positions, for example.

[0205] In various embodiments, further to the above, anvil 1130 can include one or more apertures, slots, or recesses 1179 (FIG. 50) which can be configured to receive at least a portion of housing 1170 when anvil 1130 is brought into close opposition to staple cartridge 1150, for example. In at least one embodiment, sufficient clearance can be present between housing 1170 and recess 1179 such that anvil 1130 and staple cartridge 1150 can be moved relative to each other without interference, or at least substantial interference, therebetween. In embodiments having more than one cutting member housing as outlined above, an opposing anvil can have more than one corresponding aperture for receiving the housings. In various embodiments, an anvil can include a movable cutting member and at least one housing for at least partially covering, enclosing, and/or surrounding the cutting member. In certain embodiments, although not illustrated, both an anvil and a staple cartridge can comprise at least one movable cutting member and/or at least one housing configured to at least partially cover, surround, or enclose the cutting member when they are in a proximal position, for example.

[0206] As outlined above, pusher bar assembly 1200 can be advanced distally in order to move staple sled assembly 1160 within staple cartridge assembly 1150. In various embodiments, as also outlined above, the wedge-like cam surfaces 1167 of staple sled 1162 can be moved into engagement with the sloped surfaces 1169 on staple drivers 1168 to sequentially, and/or simultaneously, drive staples from staple cartridge 1150 against anvil 1130 and form the staples into any suitable configuration, such as B-shaped configurations, for example. In at least one such embodiment, referring to FIG. 50, anvil 1130 can include one or more staple forming surfaces, such as staple pockets 1132, for example, which can be configured to deform the staples. In certain embodiments, anvil 1130 can further include a slot, channel, or groove 1133 which can be configured to slidably receive at least a portion of staple sled 1162, cutting member 1164, and/or pusher bar 1202, for example. In at least one embodiment, although not illustrated, an anvil can include an anvil plate which can be securely and/or immovably positioned within an anvil channel defined within the anvil. In various other embodiments, as illustrated in FIGS. 51 and 52 and described in detail below, anvil 1130 can include an anvil plate 1134 movably positioned within anvil channel 1136. In certain embodiments, anvil channel 1136 can include opposite side walls 1137 and, in addition, a base 1138 extending between side walls 1137. In at least one embodiment, anvil 1130 can further include a distal nose portion 1139, for example, assembled thereto wherein nose portion 1139 can be configured to be press-fit and/or snap-fit into anvil channel 1136, for example, such that nose portion 1139 can be securely retained therein. In certain embodiments, nose portion 1139 can be comprised of a soft and/or pliable material, such as rubber, for example, and can comprise any suitable shape which can facilitate the insertion of anvil 1130 into a surgical site, for example. In some embodiments, referring to FIG. 51, a nose portion, such as nose portion 1139, can be retained by an anvil by one or more listeners 1139a/b. Similarly, referring to FIG. 34, a staple cartridge channel and/or staple cartridge, such as staple cartridge 1150, for example, can include a nose portion, such as nose portion 1153, for example, which can facilitate the insertion of staple cartridge 1150 into a surgical site, for example.

[0207] As indicated above, staples can be deployed from a staple cartridge and deformed against an anvil. In various circumstances, the distance between the staple forming surfaces on anvil 1130 and staple sled 1162 can determine the amount in which the staples are deformed. For example, if the distance between anvil pockets 1132 on anvil 1130 and top surfaces 1135 on staple sled 1162 (FIGS. 43-45) is relatively large, the staples will be deformed a lesser amount as compared to when the distance between anvil pockets 1132 and sled surfaces 1135 is relatively small. Correspondingly, if the distance between anvil pockets 1132 and sled surfaces 1135 is relatively small, the staples will be deformed a greater amount as compared to when the distance between anvil pockets 1132 and sled surfaces 1135 is relatively large. Often, the distance between anvil pockets 1132 and sled surfaces 1135 is referred to as the forming height of the staples. Sometimes the forming height of the staples can be measured between the top surface, or deck, of the staple cartridge and the staple forming surfaces on the anvil. For the purpose of this application, however, any reference to a staple forming height, or the like, can include one or both manners of measurement, where appropriate, and/or any other suitable manner of measurement. In any event, as described in greater detail below, a surgical stapling instrument, such as stapling instrument 1100, for example, can include means for adjusting the staple forming height.

[0208] In various embodiments, further to the above, an anvil can include one or more forming surfaces which can be moved toward and/or away from a staple cartridge in order to set the forming height of the staples. In at least one embodiment, referring to FIGS. 50-56, anvil 1130 can include anvil plate 1134 which can be movably and/or slidably positioned within anvil channel 1136. In certain embodiments, anvil 1130 can further include one or more retention, or guide, pins 1140, wherein anvil plate 1134 can include one or more retention, or guide, slots 1141 configured to slidably receive at least a portion of pins 1140. In at least one such embodiment, pins 1140 and/or slots 1141 can be configured to define a predetermined path along which anvil plate 1134 can be moved. Referring to FIG. 51, pins 1140 and slots 1141 can be structured and arranged such that anvil plate 1134 can be moved along a linear, or at least substantially linear, path, wherein the linear path can be at least partially defined by axes 1142 and 1143, for example. Other embodiments are envisioned in which an anvil plate can be moved along a non-linear path, such as a curved and/or curvi-linear path, for example. In certain embodiments, at least a portion of pins 1140 can be retained within apertures 1144 in side walls 1137 wherein, in at least one embodiment, pins 1140 can be press-fit within apertures 1144. In any event, as described herein, pins 1140 can guide anvil plate 1134 as it is moved toward and/or away from staple cartridge 1150, for example.

[0209] In various embodiments, further to the above, a surgical stapling instrument, such as stapling instrument 1100, for example, can include one or more adjustment members configured to position a portion of an anvil, such as anvil plate 1134, for example, relative to other portions of an anvil assembly and/or an opposing staple cartridge. In certain embodiments, referring to FIGS. 51 and 52, stapling instrument 1100 can include anvil plate adjustment member 1230 which can be configured to limit the range of motion of anvil
plate 1134. In at least one such embodiment, referring to FIGS. 120 and 121, adjusting member 1230 can be positioned intermediate anvil plate 1134 in a first position in which first surface, or step, 1231 of adjusting member 1230 is positioned intermediate base 1138 of anvil channel 1136 and first positioning surface 1145 on anvil plate 1134. In such a first position, first step 1231 can define the amount of relative movement possible, or permitted, between anvil plate 1134 and anvil channel 1136. For example, when anvil 1130 is clamped against tissue as described above, anvil plate 1134 can contact the tissue and slide upwardly toward base 1138 until first positioning surface 1145 contacts first step 1231. Once surface 1145 and step 1231 are in contact, adjusting member 1230 can prevent, or at least inhibit, anvil plate 1134 from being further upward toward base 1138. In at least one embodiment, as a result, adjusting member 1230 can act as a stop such that the distance between base 1138 and tissue-contacting surface 1148 on anvil plate 1134 can be defined by a first distance 1234. While base 1138 is used as a reference datum in the present example, other portions of anvil 1130 and/or an opposing staple cartridge, for example, could be used as reference datums. When adjusting member 1230 is in its first position, as described above, second surface, or step, 1232 of adjusting member 1230 can be positioned intermediate base 1138 and second positioning surface 1146 on anvil plate 1134, and, in addition, third surface, or step, 1233 can be positioned intermediate base 1138 and third positioning surface 1147. Referring to FIG. 55, adjustment member 1230 can include two or more sets of steps, 1231, 1232, and/or 1233 and anvil plate 1134 can include two or more sets of positioning surfaces 1145, 1146, and/or 1147. While first step 1231 and first positioning surface 1145 are described above as being configured to control the position of anvil plate 1134, the second and third steps (1232, 1233) of adjustment member 1230 and the second and third positioning surfaces (1146, 1147) of anvil plate 1134, respectively, can also be configured to control the position of anvil plate 1134. For the sake of brevity, though, the present example will be described in reference to the first surface, or step 1231, as being the surface which controls the position of anvil plate 1134, although the reader will understand that the steps 1232 and 1233 can control the position of anvil plate 1134 as well.

In certain embodiments, the first position of adjustment member 1230 can provide for a relatively small, or short, staple forming height. In other embodiments, although not illustrated, the first position of an adjustment member can provide for an intermediate, a relatively large, and/or any other suitable staple forming height. In the event that the forming height associated with the first position of the adjustment member is suitable, a surgeon can proceed to use the surgical stapling instrument to staple and/or incise tissue as described above. In the event, however, that the staple forming height is unsuitable, a surgeon, or other clinician, can move adjustment member 1230 such that adjustment member 1230 can permit anvil plate 1134 to slide upwardly a different distance when anvil plate 1134 contacts tissue positioned intermediate anvil 1130 and staple cartridge 1150. In at least one such circumstance, the distance in which anvil plate 1134 is permitted to slide upwardly can be larger, thereby providing a larger forming height for the staples. Correspondingly, in other circumstances, the adjustment member can be moved such that anvil plate 1134 can slide upwardly a shorter distance when anvil plate 1134 contacts the tissue, for example, thereby providing a shorter staple forming height. While the term “upward”, and the like, can mean vertically upward, the term is not so limited; rather, “upward” can mean any direction which is toward the base of the anvil and/or away from a staple cartridge, for example. In any event, adjustment member 1230 can be moved between its first position, illustrated in FIG. 54, and a second position, illustrated in FIG. 55, in order to increase the staple forming height. As indicated by arrow “P” in FIG. 55, adjustment member 1230 can be slid proximally in order to move adjustment member 1230 between its first and second positions, although embodiments are envisioned where an adjustment member can be slid distally and/or any other suitable direction in order to adjust adjustment member 1230. Once adjustment member 1230 has been moved into its second position, referring to FIG. 55, first surface, or step, 1231 can be positioned intermediate base 1138 and second positioning surface 1146 of anvil plate 1134. In such a second position, first step 1231 can once again define the amount of relative movement permitted between anvil plate 1134 and anvil channel 1136. In at least one embodiment, similar to the above, adjusting member 1230 can act as a stop such that the distance between base 1138 and tissue-contacting surface 1148 on anvil plate 1134 can be defined by a second distance 1235.

Further to the above, adjustment member 1230 can be moved between its second position, illustrated in FIG. 55, and a third position, illustrated in FIG. 56, in order to once again increase the staple forming height. As indicated by arrow “P” in FIG. 56, adjustment member 1230 can be slid proximally in order to move adjustment member 1230 between its second and third positions. Once adjustment member 1230 has been moved into its third position, referring to FIG. 56, first surface, or step, 1231 can be positioned intermediate base 1138 and third positioning surface 1147. In such a third position, first step 1231 can once again define the amount of relative movement between anvil plate 1134 and anvil channel 1136. In at least one embodiment, similar to the above, adjusting member 1230 can act as a stop such that the distance between base 1138 and tissue-contacting surface 1148 on anvil plate 1134 can be defined by a third distance 1236. While adjustment member 1230 can be selectively moved between three positions as described above to provide three different staple forming heights, other embodiments are envisioned which comprise an adjustment member which can be moved between more than three positions to provide more than three different staple forming heights. For example, an adjustment member can be movable between four positions in order to provide four staple forming heights. Further embodiments are envisioned which comprise an adjustment member which can be moved between two positions to provide two staple forming heights. Furthermore, while surfaces, or steps, 1231, 1232, and 1233 of adjustment member 1230 are arranged in a descending order, other arrangements are envisioned in which the surfaces, or steps, are arranged in an ascending order. Other arrangements are envisioned in which the surfaces, or steps, are not necessarily arranged in either an ascending or a descending order. Similarly, positioning surfaces 1145, 1146, and 1147 of anvil plate 1134 can be arranged in an ascending order, a descending order (FIG. 53), and/or any other suitable order. Furthermore, while adjustment member 1230 can be slid along an axis, other embodiments are envisioned where an adjustment member can be moved along any suitable path such as curved and/or curvilinear paths, for example.
[0212] As described above, referring to FIG. 54, adjustment member 1230 can comprise three surfaces, or steps, 1231, 1232, and 1233 while anvil plate 1134 can comprise three corresponding adjustment surfaces 1145, 1146, and 1147. When adjustment member 1230 is in its first position, for example, first surface 1231 can be positioned such that it abuts or is adjacent to first adjustment surface 1145, second surface 1232 can be positioned such that it abuts or is adjacent to second adjustment surface 1146, and third surface 1233 can be positioned such that it abuts or is adjacent to third adjustment surface 1147. As adjustment member 1230 is slid relative to anvil plate 1134, as described above and referring to FIGS. 55 and 56, surfaces 1231, 1232, and 1233 of adjustment member 1230 can be sequentially indexed relative to surfaces 1145, 1146, and 1147 of anvil plate 1134. In at least one such embodiment, an adjustment member can have the same number of steps as the number of positioning surfaces on an anvil plate. Other embodiments are envisioned wherein an adjustment member has more steps than positioning surfaces on the anvil plate. In at least one such embodiment, an anvil plate can include one positioning surface wherein the steps of an adjustment member can be selectively utilized to limit the upward movement of the anvil plate, for example. In various embodiments, referring generally to adjustment member 1230 and anvil plate 1134, an anvil plate may include one positioning surface, such as positioning surface 1145, for example, where steps 1231, 1232, and 1233 of adjustment member 1230, for example, can be selectively positioned intermediate base 1138 and positioning surface 1145. In such embodiments, first step 1231 can have a first thickness or height which can stop, or limit, the upward movement of anvil plate 1134 so as to define a first staple forming height, second step 1232 can have a second thickness or height which can stop, or limit, the upward movement of anvil plate 1134 so as to define a second staple forming height, and, in addition, third step 1233 can have a third thickness or height which can stop, or limit, the upward movement of anvil plate 1134 so as to define a third staple forming height. In at least one embodiment, the thickness or height of steps 1231, 1232, and/or 1233 can be measured between a back surface 1237 of adjustment member 1230 and a surface on the steps (1231, 1232, 1233) which will contact anvil plate 1134. In various embodiments, the difference in height, or thickness, between first step 1231 and second step 1232 can be the same, or at least substantially the same, as the difference in height, or thickness, between second step 1232 and third step 1233. In at least one such embodiment, as a result, the step heights can increase at a linear rate, or an at least substantially linear rate. In alternative embodiments, the difference in height, or thickness, between the first and second steps can be different than the difference in height, or thickness, between the second and the third steps. In at least one such embodiment, the first, second, and third steps may not increase or decrease in height, or thickness, at a linear rate; rather, although not illustrated, the steps may increase or decrease in height, or thickness, in a non-linear and/or geometric rate.

[0213] As described above, an adjustment member, such as adjustment member 1230, for example, can be movable between two or more positions. In various embodiments, a surgical stapling instrument can include an actuator configured to move the adjustment member. In at least one embodiment, referring to FIGS. 50-53, surgical stapling instrument 1100 can include actuator 1250 which can be operably attached to adjustment member 1230 such that a force can be applied to actuator 1250 and transmitted to adjustment member 1230. In certain embodiments, actuator 1250 can include grasping portions, or handles, 1252 which can be configured to be grasped by a surgeon, for example, in order to advance or retract adjustment member 1230 within anvil 1130 as described above. In certain embodiments, grasping portions 1252 can extend from actuator body 1251, wherein actuator body 1251 can include one or more apertures, slots, or cavities 1253 which can be configured to receive at least a portion of adjustment member 1230. In at least one such embodiment, referring to FIG. 52, adjustment member 1230 can include lock 1254 extending therefrom, wherein at least a portion of lock 1254 can be received within aperture 1253 so as to retain actuator body 1251 to adjustment member 1230. In at least one embodiment, legs 1255 which can be deflected when they are inserted into aperture 1253 but resistently return, or at least partially return, to their unflexed position after feet 1256 of legs 1255 are sufficiently pushed through aperture 1253. In at least one such embodiment, feet 1256 can prevent, or at least inhibit, actuator body 1251 from being detached from adjustment member 1230.

[0214] In various embodiments, further to the above, surgical stapling instrument 1100 can further include a detent mechanism which can be configured to hold, or releasably hold, actuator 1250 and/or adjustment member 1230 in position. In at least one embodiment, referring to FIG. 52, detent member 1260 can be attached to actuator 1250 wherein, in at least some embodiments, actuator body 1251 can include one or more channels, grooves, or recesses 1257 which can be configured to receive and/or retain a detent body 1261 of detent member 1260 therein. In at least one embodiment, detent body 1261 can include one or more apertures 1263, and/or any other suitable channels, slots, or grooves, which can be configured to receive one or more fasteners for securing detent body 1261 to actuator 1251, for example. Detent member 1260 can further include detent legs 1262 which can be configured to engage one or more recesses, apertures, or grooves 1101 (FIGS. 35-40) in first frame portion 1110, for example. More particularly, referring to FIGS. 34 and 35, each side flange 1128 can include one or more recesses 1101 (1101a, 1101b, and 1101c) defined therein wherein detent legs 1262 can be biased into engagement with the top surfaces of side flanges 1128 such that detent legs 1262 can be slid into, and slid out of, recesses 1101. In the illustrated embodiment, each side flange can include three recesses 1101 which can be configured to removably hold actuator 1250 in a first, distal position, a second, lock 1254 in position, and a third, proximal position, wherein the first, second, and third positions of actuator 1250 can respectively correspond with the first, second, and third positions of adjustment member 1230 described above. For example, when actuator 1250 is in its first, distal position, detent legs 1262 of detent member 1260 can be positioned within recess 1101a so as to removably retain actuator 1250 and adjustment member 1230 in their first positions. Upon the application of a sufficient force, actuator 1250 can be moved proximally into its second position such that detent legs 1162 are positioned within recess 1101b and actuator 1250 and adjustment member 1230 are retained in their second positions. Similarly, upon the application of a sufficient force, actuator 1250 can be moved proximally into its third position such that detent legs 1162 are positioned within recess 1101c and actuator 1250 and adjustment member 1230 are retained in their third positions.
In various embodiments, detent legs 1162 can be configured such that actuator 1250 can be returned to its first and/or second positions.

[0215] As described above, adjustment member 1230 can be moved along a pre-determined path between two or more positions by actuator 1250. In various embodiments, surgical stapling instrument 1100, for example, can include one or more guides for controlling or limiting the movement of adjustment member 1230 and/or actuator 1250. In some embodiments, adjustment member 1230 can be closely received between side walls 1137 of anvil 1130 such that side walls 1137 can guide adjustment member 1230. In at least one such embodiment, side walls 1137 can be configured to control or limit the lateral or side-to-side movement of adjustment member 1230. In various embodiments, detent leg 1162 of detent member 1160 can comprise resilient members which can be configured to apply an upward biasing or pulling force on adjustment member 1230 so as to position adjustment member 1230 against, or at least adjacent to, base 1138 and intermediate side walls 1137. In certain embodiments, referring to FIG. 52, base 138 of anvil 1130 can further include guide slot 1149 which can be configured to receive at least a portion of adjustment member 1230 and/or actuator 1250 therein such that guide slot 1149 can limit the movement of adjustment member 1230 and actuator 1250. In at least one such embodiment, lock 1254 of adjustment member 1230 can be configured to extend through guide slot 1149 such that, when lock 1254 is inserted into aperture 1253 of actuator 1250 as described above, base 1138 of anvil 1130 can be captured intermediate adjustment member 1230 and actuator 1250. In certain embodiments, guide slot 1149 can be configured to limit the movement of lock 1254 such that adjustment member 1230 can be prevented, or at least inhibited, from being moved distally when adjustment member 1230 is in its first, or distal-most, position and/or, similarly, prevented, or at least inhibited, from being moved proximally when adjustment member 1230 is in its third, or proximal-most, position.

[0216] In various embodiments, further to the above, a detent member, similar to detent member 1260, for example, can be utilized to bias first handle portion 1102 and second handle portion 1104 away from one another. In at least one embodiment, referring to FIG. 70, surgical stapling instrument 1100 can include a detent member 1260 configured to position first handle portion 1102 and second handle portion 1104 such that a gap exists between anvil 1130 and staple cartridge 1150. Such a feature, as outlined above, can allow a surgeon to easily manipulate the stapler without having to hold the first and second handle portions apart from one another. In certain embodiments, detent member 1260 can be sufficiently mounted to second handle portion 1104 such that detent legs 1262 extending from detent member 1260 can contact flanges 1128 and, when compressed, apply a biasing force to the first and second handle portions. As seen in FIG. 70, legs 1262 can contact surfaces 1101d on flanges 1128. In order to compress detent legs 1262, latch mechanism 1180 can be moved into a partially-closed position such that latch arms 1188 can engage, and at least partially surround, latch projections 1131. In this configuration, a surgeon can manipulate the instrument and, when satisfied with its position, move latch mechanism 1180 into a closed position and further compress detent legs 1262. Similar to the above, detent member 1260 can be affixed, or otherwise operably engaged with, actuator 1250 such that, when actuator 1250 is moved between its first, second, and third positions as described above, legs 1262 can engage recesses 1101a, 1101b, and 1101c, respectively. In at least one such embodiment, as a result, actuator 1250 can have a pre-staged position in which actuator 1250 is positioned distally with respect to its first position and, in addition, surfaces 1101d can comprise pre-stage surfaces against which legs 1262 can be positioned when actuator 1250 is in its pre-staged position.

[0217] As outlined above, an adjustment member can be slid, or translated, between first and second positions so as to adjust the forming height of staples deployed by a surgical stapling instrument. In various embodiments, although not illustrated, an adjustment member can be configured to positively displace an anvil plate toward and/or away from an opposing staple cartridge, for example. In at least one such embodiment, a surgical stapling instrument can include one or more biasing members, such as springs, for example, configured to position the anvil plate against the adjustment member such that, when the adjustment member is moved between its first and second positions, the adjustment member can displace the anvil plate between first and second positions in order to set first and second staple forming heights. In various embodiments, as a result of the above, an adjustment member can be configured to cam a portion of an anvil into position. In at least one such embodiment, an adjustment member can be slid along an axis in order to positively displace an anvil plate. In other embodiments, a rotatable adjustment member can be configured to positively displace an anvil plate toward and/or away from a staple cartridge, for example.

[0218] Further to the above, as described in greater detail below, an adjustment member can be rotated to adjust the staple forming height. Referring to FIGS. 57-69, surgical instrument 1100 can include, similar to the above, a first handle portion 1102, a second handle portion 1104, and a latching mechanism 1180 which can be utilized to clamp tissue intermediate anvil 1130 and staple cartridge 1150. Referring to FIG. 58, also similar to the above, latching mechanism 1180 can be pivotably coupled to first portion 1102 by one or more pivot pins 1182, wherein latching mechanism 1180 can include one or more latch arms 1188 which can be configured to engage second portion 1104 and latch the first and second handle portions together. Also similar to the above, referring to FIGS. 58 and 60, surgical instrument 1100 can further include pusher bar assembly 1200 which can be configured to advance a cutting member and/or staple sled within end-effector 1120. In at least one such embodiment, pusher bar assembly 1200 can include a proximal end 1203 and an actuator 1204, wherein actuator 1204 can be rotatably mounted to proximal end 1203 and selectively positioned on first and second sides of stapling instrument 1100. In various embodiments, surgical stapling instrument 1100 can comprise the same, or similar, features to those described in connection with surgical stapling instrument 1100 and can be operated in the same manner, or a similar manner, as instrument 1100 and, as a result, such details are not repeated herein.

[0219] In various embodiments, referring to FIG. 60, surgical instrument 1100 can include a rotatable adjustment member 1230 which can be selectively positioned in at least first and second positions so as to provide different staple forming heights. In certain embodiments, surgical instrument 1100 can include an actuator 1250 which can be operably connected to adjustment member 1230 such that actuator
1250' can move adjustment member 1230' between at least its first and second positions. In at least one embodiment, referring to FIG. 61, actuator 1250' can include actuator body 1251' and grasping portion, or handle, 1252'. Actuator body 1251' can include an aperture 1258' which can be configured to receive a proximal end 1238' of adjustment member 1230'. Such that rotational motion, torque, and/or forces can be transmitted between actuator body 1250' and adjustment member 1230' In at least one such embodiment, referring to FIG. 69, aperture 1258' can comprise a non-circular profile and/or a profile which includes one or more flat drive surfaces configured to transmit rotational motion between actuator body 1251' and actuator 1230'. In certain embodiments, aperture 1258' can be sized and configured to closely receive proximal end 1238' of actuator 1230'. In at least one embodiment, aperture 1258' can be configured to receive proximal end 1238' in a press-fit and/or snap-fit arrangement. In various embodiments, referring again to FIG. 61, handle portion 1104' can include one or more slots 1259' which can be configured to permit at least a portion of actuator body 1251' to extend therethrough such that grasping portion 1252' can be assembled to actuator body 1251' with at least a portion of handle portion 1104' positioned therethrough. In at least one such embodiment, second handle portion 1104' can further include recess 1253' which can be configured such that at least a portion, if not all, of grasping portion 1252' is positioned within recess 1253'. In certain embodiments, recess 1253' can be configured such that grasping portion 1252' does not extend above the top surface of second handle portion 1104' although, in other embodiments, an upper portion of grasping portion 1252' can extend above second handle portion 1104', as illustrated in FIG. 63, such that grasping portion 1252' can be easily accessed by a surgeon.

[0220] In various embodiments, as outlined above, an adjustment member can be rotatable between at least first and second positions in order to adjust the forming height of staples deployed by a surgical stapler. In certain embodiments, referring to FIG. 61, a surgical stapling instrument can include an adjustment member rotatably positioned within an anvil wherein the adjustment member can be configured to limit the relative movement of a movable anvil portion. In at least one such embodiment, surgical stapling instrument 1100' can include an anvil plate 1134' which can be slidably retained within anvil channel 1136' by retention, or guide, pins 1140', wherein guide pins 1140' can be configured to allow anvil plate 1134' to slide upwards when anvil plate 1134' comes into contact with tissue as described above. Referring to FIGS. 60, 63, and 64, adjustment member 1230' can be positionable in a first position, or orientation, such that it can limit the upward movement of anvil plate 1134' within anvil channel 1136' and dictate the staple forming height of the staples. In at least one such embodiment, referring to FIGS. 63 and 64, adjustment member 1230' can include opposing first surfaces 1231' which can be positioned intermediate base 1138' of anvil channel 1136' and positioning surface 1145' of anvil plate 1134' such that, when positioning surface 1145' contacts one of first surfaces 1231', tissue-contacting surface 1148' of anvil plate 1134' can be positioned a first distance 1234' away from a datum surface 1129' on anvil 1130', for example. Correspondingly, forming surfaces 1132' can be positioned a first distance away from a staple cartridge such that, when staples are deployed from the staple cartridge, the staples can be deformed to a first staple height. Further to the above, a first diameter 1241' can be defined between first surfaces 1231' wherein the first diameter 1241' can define the maximum upward position of anvil plate 1134' within anvil channel 1136'.

[0221] As indicated above, adjustment member 1230' can be rotated in order to adjust the forming height of the staples. In various embodiments, adjustment member 1230' can be rotated between its first position, or orientation, (FIGS. 63 and 64) and a second position, or orientation (FIGS. 65 and 66). In at least one embodiment, referring to FIGS. 65 and 66, handle 1252' can be rotated in a direction indicated by arrow “A” in order to move adjustment member 1230' between its first and second positions. Similar to the above, when actuator 1230' is in its second position, or orientation, actuator 1230' can limit the upward movement of anvil plate 1134' within anvil channel 1136' and dictate the staple forming height of the staples. In at least one such embodiment, referring to FIGS. 65 and 66, adjustment member 1230' can include opposing second surfaces 1232' which can be positioned intermediate base 1138' and positioning surface 1145' such that, when positioning surface 1145' contacts one of second surfaces 1232', tissue-contacting surface 1148' of anvil plate 1134' can be positioned a second distance 1235' away from datum surface 1129', for example. Correspondingly, forming surfaces 1132' can be positioned a second distance away from a staple cartridge such that, when staples are deployed from the staple cartridge, the staples can be deformed to a second staple height. In various embodiments, similar to the above, a second diameter 1242' can be defined between second surfaces 1232', wherein second diameter 1242' can define the maximum upward position of anvil plate 1134' within anvil channel 1136'. Although first surfaces 1231' and second surfaces 1232' can be defined by flat, or at least substantially flat, surfaces, other embodiments are envisioned in which the first and second surfaces 1231' and 1232' can include at least partially arcuate, or curved, contours. In any event, referring to FIG. 60, adjustment member 1230' may include one or more clearance slots 1240' which can be configured to provide clearance between actuator 1230' and retention pins 1140'. Clearance slots 1240' can be configured to provide clearance between actuator 1230' and retention pins 1140' when actuator 1230' is in its first position, second position, and/or any other suitable position.

[0222] In various embodiments, further to the above, adjustment member 1230' can be rotated between its first position, or orientation, (FIGS. 63 and 64) and a third position, or orientation (FIGS. 67 and 68). In at least one embodiment, referring to FIGS. 67 and 68, handle 1252' can be rotated in a direction indicated by arrow “B” in order to move adjustment member 1230' between its first and third positions. Similar to the above, when actuator 1230' is in its third position, or orientation, actuator 1230' can limit the upward movement of anvil plate 1134' within anvil channel 1136' and dictate the staple forming height of the staples. In at least one such embodiment, referring to FIGS. 67 and 68, adjustment member 1230' can include opposing third surfaces 1233' which can be positioned intermediate base 1138' and positioning surface 1145' such that, when positioning surface 1145' contacts one of third surfaces 1233', tissue-contacting surface 1148' of anvil plate 1134' can be positioned a third distance 1236' away from datum surface 1129', for example. Correspondingly, forming surfaces 1132' can be positioned a third distance away from a staple cartridge such that, when staples are deployed from the staple cartridge, the staples can be deformed to a third staple height. In various embodiments,
similar to the above, a third diameter 1243' can be defined between third surfaces 1233', wherein third diameter 1243' can define the maximum upward position of anvil plate 1134' within anvil channel 1136'. Referring once again to FIGS. 67 and 68, third surfaces 1233' can be defined by an at least partially arcuate contour, although other embodiments are envisioned in which third surfaces 1233' can include flat, or at least substantially flat, contours. In at least one embodiment, adjustment member 1230' can be configured such that the largest distance, or diameter, between the arcuate third surfaces 1233' can be utilized to define the third staple height.

[0223] As described above, referring to FIGS. 63 and 64, adjustment member 1230' can be positioned in a first position, or orientation, to set a first forming height for the staples deployed by surgical stapling instrument 1100'. As also described above, referring to FIGS. 65 and 66, actuator 1250' can be utilized to move adjustment member 1230' into its second position, or orientation, to set a second forming height for the staples. To do this, in at least one embodiment, a force can be applied to handle 1252' which can cause handle 1252', and adjustment member 1230' attached thereto, to rotate in a direction indicated by arrow "A". In at least one embodiment, adjustment member 1230' and/or actuator 1250' can be sufficiently retained such that, when adjustment member 1230' is rotated, adjustment member 1230' can be rotated about an axis, such as axis 1245' (FIG. 60), for example. In at least one embodiment, referring to FIG. 58, the proximal end 1203' of pusher bar assembly 1200' can include one or more grooves, channels, or recesses 1201' which can be configured to receive and/or retain at least a portion of adjustment member 1230' and/or actuator 1250' therein. In any event, as illustrated in FIGS. 63-66, the second position, or orientation, of adjustment member 1230' can allow anvil plate 1134' to slide a larger distance within anvil channel 1136' as compared to when adjustment member 1230' is in its first position. In at least one embodiment, as a result, the second staple forming height can be larger than the first staple forming height. As also described above, referring to FIGS. 67 and 68, actuator 1250' can be utilized to move adjustment member 1230' into its third position, or orientation, to set a third forming height for the staples. To do this, in at least one embodiment, a force can be applied to handle 1252' which can cause handle 1252', and adjustment member 1230' attached thereto, to rotate in a direction indicated by arrow “B”. As illustrated in FIGS. 63, 64, 67, and 68, the third position, or orientation, of adjustment member 1230' can allow anvil plate 1134' to slide a smaller distance within anvil channel 1136' as compared to when adjustment member 1230' is in its first position. In at least one embodiment, as a result, the first and second staple forming heights can be larger than the third staple forming height. In at least one such embodiment, the first position of adjustment member 1230', and actuator 1250', can represent an intermediate position, wherein adjustment member 1230' can be selectively moved into its second and third positions directly from its first position. In effect, the first position of adjustment member 1230' can represent an intermediate staple height, wherein the second and third staple positions of adjustment member 1230' can represent taller and shorter staple heights, respectively. In certain embodiments, referring to FIG. 57, surgical stapling instrument 1100' can include one or more indicia thereon which can be configured to convey the staple forming heights, or at least relative forming heights, that can be selected. For example, second handle portion 1104' can include a first indicium 1245' which can indicate an intermediate, or first, staple height, a second indicium 1246' which can indicate a taller, or second, staple height, and, in addition, a third indicium 1247' which can indicate a shorter, or third, staple height.

[0224] In various embodiments, further to the above, one or more of first surfaces 1231', second surfaces 1232', and third surfaces 1233' can comprise or define, or at least partially comprise or define, a perimeter, or circumference, of adjustment member 1230'. As discussed above, owing to the first, second, and third diameters (1241', 1242', and 1243') defined by the first, second, and third surfaces (1231', 1232', and 1233'), respectively, the perimeter, or circumference, of adjustment member 1230' may be non-circular. In certain embodiments, though, the perimeter, or circumference of adjustment member 1230', may be symmetrical, substantially symmetrical, and/or non-symmetrical. In various embodiments, further to the above, an adjustment member can comprise a cam rotatably positioned intermediate base 1138' of anvil 1130' and adjustment surface 1145' of anvil plate 1134', for example. In at least one such embodiment, one or more of first surfaces 1231', second surfaces 1232', and third surfaces 1233', for example, can comprise or define a cam profile which, similar to the above, can be configured to either positively position anvil plate 1134' and/or provide a stop against which anvil plate 1134' can be positioned. In any event, although not illustrated, various embodiments are envisioned in which an adjustment member can be slid and rotated in order to set two or more staple forming heights for staples deployed by a surgical stapling instrument. In at least one such embodiment, an adjustment member can comprise a cam profile which can be defined along the length of the adjustment member wherein longitudinal and/or rotational movement can be utilized to move the cam profile between at least first and second positions.

[0225] In various embodiments, similar to the above, surgical instrument 1100' can further include a detent mechanism configured to hold, or at least releasably hold, actuator 1250' in position. In at least one embodiment, referring to FIGS. 58 and 59, surgical instrument 1100' can further include detent member 1260' comprising detent body 1261' and one or more detent legs 1262'. Referring to FIG. 59, detent body 1261' can include one or more grooves, recesses, or channels 1263' which can be configured to receive at least a portion of proximal end 1105' of second handle portion 1104' therein such that detent member 1260' can be retained in position. In at least one such embodiment, proximal end 1105' can further include one or more grooves, channels, or recesses 1265' which can be configured to closely receive detent member 1260'. In certain embodiments, at least a portion of detent body 1261', such as channel 1263', for example, can be press-fit, snap-fit, and/or otherwise suitably retained in recess 1265'. As also illustrated in FIG. 59, each detent leg 1262' of detent member 1260' can include one or more projections 1264' extending therefrom which can be configured to engage actuator body 1251' and releasably hold actuator 1250' in position. In at least one embodiment, referring to FIG. 60, actuator body 1251' can include one or more recesses, or holes, 1269' which can be configured to receive a projection 1264'. When a projection 1264' is positioned within recess 1269', the projection can be configured to hold actuator 1250' in its first position, for example, until a sufficient force is applied to actuator 1250' so as to cause the projection 1264' to be displaced out of recess 1269'. More particularly, the force applied to actuator 1250' can trans-
mitted to the projection 1264' and, owing to cooperating surfaces between the projection 1264' and recess 1269', the detent leg 1262' associated with the projection 1264' can be flexed or moved proximally to allow actuator body 1251' to be moved relative thereto. In order to accommodate such proximal movement, referring to FIG. 58, recess 1265' can include elongate portions 1266' which can each be configured to receive at least a portion of legs 1262' such that legs 1262' can move relative to handle portion 1104'. As actuator 1250' is moved into either its second or third position, actuator body 1251' can contact a projection 1264' extending from another leg 1262' and deflect the leg 1262' proximally such that, once actuator 1250' is in its second or third positions, the leg 1262' can spring forward, or distally, such that the projection 1264' can be secured within recess 1269'. In at least one embodiment, further to the above, the interaction between projections 1264' and the sidewalks of recesses 1269' can be such that actuator 1250' can be securely held in one of its first, second, and third positions, for example, yet permit actuator 1250' to be moved upon a sufficient application of force. In such embodiments, the detent member 1260' can prevent, or at least inhibit, actuator 1250' and, correspondingly, adjustment member 1230' from being unintentionally displaced.

[0226] As discussed above and as shown in FIG. 35, each side flange 1128 of first handle portion 1102 can include a notch, or recess, 1127, for example, which can be configured to receive one or more latch projections 1131, for example, extending from anvil 1130, and/or any other suitable portion of second handle portion 1104. As also discussed above, referring primarily to FIGS. 35 and 36, first handle portion 1102 can further include latching mechanism 1180 rotatably mounted thereto which can be utilized to engage latch projections 1131 extending from second handle portion 1104 and secure the first and second handle portions 1102, 1104 together. Latching mechanism 1180 can include one or more latch arms 1188 extending therefrom which can be configured to engage latch projections 1131 and pull and/or secure projections 1131 within recesses 1127 as illustrated in FIG. 40. Referring to FIG. 39, at least one of latch arms 1188 can include a distal hook 1189 which can be configured to wrap around at least a portion of projections 1131 so as to encompass or surround, or at least partially encompass or surround projections 1131. In at least one embodiment, latch arms 1188 can act as an over-center latch to maintain latching mechanism 1180 in its latched, or closed, position.

[0227] In various embodiments, referring now to FIG. 71, each projection 1131 can comprise a slot, or groove, 1190 positioned intermediate side wall 1191 and an enlarged end, or head, 1192 of projection 1131, wherein the slot 1190 can be configured to receive at least a portion of latch arm 1188. More particularly, in at least one embodiment, the slot 1190 can have a width which is greater than the width of the latch arm 1188 such that, when the latch arm 1188 is engaged with the projection 1131, the latch arm 1188 can enter into slot 1190. In some circumstances, the width of each slot 1190 may be slightly larger than the width of a latch arm 1188 such that the latch arm is closely received within the slot 1190. In various circumstances, the slot 1190, the sidewall 1191, and the head 1192 of projection 1131 can be sized and configured so as to prevent, or at least limit, relative lateral movement, i.e., movement away from or to the sides of anvil 1130, between latch arms 1188 and projection 1131. Further to the above, however, the latch arms 1188 can slide longitudinally within the grooves 1190 as the latch arms 1188 move the projections 1131 into the recesses 1127 in first portion 1102. Owing to such relative sliding movement between latch arms 1188 and projections 1131, frictional forces can be generated therebetween which can resist the movement of latch arms 1188. In various circumstances, the magnitude of such frictional forces can be significant when the normal, or perpendicular, contact forces between the latch arms 1188 and the sidewalks of groove 1190 are large. In many circumstances, as a result, the operator of the surgical instrument has to overcome these frictional forces when actuating clamping mechanism 1180.

[0228] In various alternative embodiments, referring now to FIGS. 72 and 73, a surgical instrument can comprise one or more latch projections having a rotatable bearing which can reduce the magnitude of the friction forces between the latch arms of a latching mechanism and the latch projections. In at least one embodiment, an anvil 1330, which can be substantially similar to anvil 1130 in many respects, can comprise a latch projection 1331 extending from each side thereof, wherein each latch projection 1331 can comprise a rotatable bearing 1393. In use, the latch arms 1188 of latching mechanism 1180, for example, can contact the rotatable bearings 1393 in order to position the latch projections 1331 in recesses 1127. In various circumstances, the latch arms 1188 can slide across the surface, or outer diameter, of bearings 1393; however, as bearings 1393 can rotate relative to the latch arms 1188, the magnitude of the frictional forces between the latch arms 1188 and projections 1331 can be lower than the magnitude of the frictional forces between latch arms 1188 and projections 1131. Owing to such lower frictional forces, a lower closing, or clamping, force may be required to actuate clamping mechanism 1180, for example.

[0229] In various embodiments, referring primarily to FIG. 74, each rotatable bearing 1393 can comprise a circular, or round, outer diameter 1394 and, in addition, a circular, or round, bearing aperture 1395 extending therethrough. In certain embodiments, each projection 1331 can further comprise a shaft portion 1396 extending from sidewall 1391 and an enlarged end, or head, 1392 extending from shaft portion 1396, wherein, as illustrated in FIG. 64, the shaft portion 1396 can extend through the bearing aperture 1395 of rotatable bearing 1393. In various embodiments, the shaft portion 1396 can comprise a circular, or round, outer diameter which can be closely received within bearing aperture 1395 such that there is little, if any, relative radial movement therebetween. The diameter of the bearing aperture 1395, however, may be sufficiently larger than the outer diameter of shaft portion 1396 such that bearing 1393 can rotate relative to shaft portion 1396 about an axis 1399. In various embodiments, the rotatable bearing 1393 can be retained on shaft portion 1396 by the enlarged head 1392. More particularly, in at least one embodiment, the enlarged head 1392 may be larger than, or define a larger diameter than, the diameter of bearing aperture 1395 such that rotatable bearing 1393 cannot slide off the end of shaft portion 1396. In certain embodiments, the sidewall 1391 and the head 1392 can define a gap distance therebetween and, in addition, the bearing 1393 can comprise a width, wherein the gap distance can be larger than the width of bearing 1393. In at least one embodiment, the gap distance may be slightly larger than the width of bearing 1393 such that bearing 1393 does not tilt, or at least substantially tilt, relative to axis 1399, for example.

[0230] As discussed above, the latch arms 1188 of latching mechanism 1180 can be configured to engage bearings 1393.
and position bearings 1393 within recesses 1127. In various alternative embodiments, referring primarily to FIG. 73, a surgical instrument can comprise a latching mechanism 1380 which can comprise first and second latch arms 1388 extending therefrom on opposite sides of anvil 1331 and staple cartridge channel 1324. In use, similar to the above, the latch arms 1388 can contact bearings 1393 in order to move bearings 1393 into recesses 1327 in staple cartridge channel 1324 and move anvil 1331 toward staple cartridge channel 1324. Such movement is illustrated with phantom lines in FIG. 74. In various embodiments, each latch arm 1388 can at least partially define a groove, or slot, 1397 therein, wherein each slot 1397 can be configured to receive a bearing 1393. In at least one embodiment, a slot 1397 can comprise a first drive surface, or sidewall, 1398a which can be positioned against bearing 1393 and, as a closing force is applied to latching mechanism 1380, the latch arm 1388 can apply a closing force to the bearing 1393. In such circumstances, the bearing 1393 can move further into slot 1397 as latching mechanism 1380 is rotated into its closed position. In various circumstances, the slot 1397 can further comprise a second drive surface, or sidewall, 1398b which can be positioned against another and/or opposite side of bearing 1393 such that an opening force can be applied to the bearing 1393 via latch arm 1388. As the latching mechanism 1380 is moved into its open position, the bearing 1393 can move out of slot 1397. In any event, the first drive surface 1398a and the second drive surface 1398b can define a slot width therebetween which can be larger than the outside diameter of bearing 1393 such that bearing 1393 can move within slot 1397. In some embodiments, the slot width may be slightly larger than the outside diameter of bearing 1393. In at least one embodiment, at least portions of the first drive surface 1398a and the second drive surface 1398b can be parallel, or at least substantially parallel, to one another. In at least one such embodiment, at least portions of the first drive surface 1398a can be positioned opposite the second drive surface 1398b.

[0231] As described above, a surgical stapling instrument can be configured to deform one or more surgical staples between a first, undeployed, configuration and a second, deployed, configuration. In various embodiments, referring now to FIG. 72, a surgical staple, such as staple 1400, for example, can comprise a base 1402, a first leg, or deformable member, 1404 extending from base 1402, and, in addition, a second leg, or deformable member, 1406 extending from base 1402. In certain embodiments, the base 1402, the first leg 1404, and the second leg 1406 can be comprised of a continuous wire, wherein, in at least one embodiment, the first leg 1404 and the second leg 1406 can each be bent in a direction which is perpendicular to the base 1402 prior to staple 1400 being inserted into and deformed by a surgical stapler. More particularly, the staple 1400 can be manufactured such that base 1402 is oriented along a baseline 1401 and such that the legs 1404 and 1406 are oriented along lines 1409 and 1411, respectively, which are perpendicular, or at least substantially perpendicular, to the baseline 1401. In various embodiments, the first leg 1404 can be positioned at a first end of base 1402 and the second end 1406 can be positioned at a second end of base 1402, wherein, in at least one embodiment, a mid-line 1403 can be defined which extends through a midpoint of base 1402 and which extends in a direction which is perpendicular to baseline 1401. The staple 1400 can be configured such that the base 1402, first leg 1404, and second leg 1406 lie, or at least substantially lie, in the same, or common, plane when the staple 1400 is in its first, or undeployed, configuration. In such embodiments, the baseline 1401, along which the base 1402 is oriented, and the perpendicular lines 1409 and 1411, along which the legs 1404 and 1406 are oriented, can lie in the same plane.

[0232] In various embodiments, further to the above, the continuous wire comprising the base 1402, the first leg 1404, and the second leg 1406 can be comprised of titanium and/or stainless steel, for example. In at least one embodiment, the first leg 1404 can comprise a first end 1405 and the second leg 1406 can comprise a second end 1407, wherein the ends 1405 and 1407 can each comprise a sharp, or chisel, tip which can be configured to puncture bone and/or tissue. In use, the staple 1400 can be deformed by a surgical stapler in order to capture tissue, for example, within the staple 1400. In various embodiments, the staple 1400 can be deployed from a staple cartridge such that the ends 1405 and 1407 of staple legs 1404 and 1406, respectively, contact an anvil positioned opposite the staple 1400. In such circumstances, a first compressive force F1 can be applied to the first leg 1404 and a second compressive force F2 can be applied to the second leg 1406 while the base 1402 is supported by at least a portion of the staple cartridge. As described in greater detail below, the anvil can comprise a staple pocket which can apply the first compressive force F1 to the first leg 1404 such that the end 1405 of staple leg 1404 is moved toward the base 1402. Similarly, the staple pocket can apply the second compressive force F2 to the second staple leg 1406 such that the end 1407 of staple leg 1404 is also moved toward base 1402. In addition to the above, as also discussed in greater detail below, referring now to FIGS. 83-85, the staple pocket can bend the first staple leg 1404 to a first side of base 1402 and the second staple leg 1406 to a second, or opposite, side of base 1402.

[0233] In various embodiments, referring to FIGS. 82 and 83, the first leg 1404 of staple 1400 can be bent such that the end 1405 of the first leg 1404 is moved toward the base 1402 and toward the second leg 1406 when the first leg 1404 is deformed by the first compressive force F1. In at least one embodiment, the end 1405 can be moved from a first side 1410 of midline 1403, as illustrated in FIG. 82, to a second side 1412 of midline 1403, as illustrated in FIG. 83. Similarly, the second leg 1406 of staple 1400 can be bent such that the end 1407 of the second leg 1406 is moved toward the base 1402 and toward the first leg 1404 when the second leg 1406 is deformed by the second compressive force F2. In at least one embodiment, the end 1407 can be moved from a second side 1410 of midline 1403, as illustrated in FIG. 82, to a first side 1410 of midline 1403, as illustrated in FIG. 83. In the deployed, or deformed, configuration of staple 1400, as illustrated in FIG. 83, the ends 1405 and 1407 of staple legs 1404 and 1406 can extend across the midline 1403 in such a way that they form an angle therebetween. More particularly, the end 1405 of the first leg 1404, when it is in its deformed configuration, can extend along or with respect to a first axis 1414 and, similarly, the end 1407 of the second leg 1406, when it is in its deformed configuration, can extend along or with respect to a second axis 1416 such that the first axis 1414 and the second axis 1416 define an angle 1417 therebetween. In some embodiments, the angle 1417 may be approximately 90 degrees, for example. In certain embodiments, the angle 1417 may be in a range between approximately 0.1 degrees and approximately 89 degrees, for example. In various embodiments, the angle 1417 may be greater than 90 degrees,
while, in at least one embodiment, the angle 1417 may be greater than approximately 90 degrees but less than 180 degrees, for example.

[0234] In various embodiments, further to the above, the first axis 1414 and the second axis 1416 can, in various embodiments, be oriented, or crossed, at a transverse angle with respect to each other, i.e., at least when the staple 1400 is viewed from the side or elevational view of FIG. 83. More particularly, upon reviewing FIG. 85, it becomes evident that, although axes 1414 and 1416 extend in transverse directions when viewed from the side (FIG. 83), the axes 1414 and 1416 may not, in at least one embodiment, actually intersect one another. In such embodiments, when viewing the staple 1400 from the top or bottom (FIG. 85), for example, the axes 1414 and 1416 may extend in parallel, or at least substantially parallel, directions. Furthermore, in various embodiments, the reader will note that the first axis 1414 and the second axis 1416 are not perpendicular with baseline 1401. Stated another way, the end 1405 of first staple leg 1404 and the end 1407 of second staple leg 1406 are not pointing directly downward toward base 1402 and baseline 1401. In at least one such embodiment, the first axis 1414 and the second axis 1416 can each extend at an acute angle with respect to baseline 1401, for example.

[0235] As described above, a surgical instrument can be configured to deform the staple 1400 of FIG. 82, for example, between an undeformed shape (FIG. 82) and a deformed shape (FIG. 83). In various embodiments, as also described above, the surgical instrument can comprise an anvil having a staple pocket configured to receive and deform at least a portion of the staple. In certain embodiments, referring now to FIG. 75, an anvil can comprise a tissue-contacting surface 1501 and a plurality of staple pockets 1500 formed therein, wherein each staple pocket 1500 can be configured to deform a staple 1400. In various embodiments, each staple pocket 1500 can comprise a longitudinal axis 1599 (FIG. 76) and, in addition, a first forming cup 1502 and a second forming cup 1504 positioned relative to the longitudinal axis 1599. In use, the first forming cup 1502 can be configured to receive the first staple leg 1404 of staple 1400 and the second forming cup 1504 can be configured to receive the second staple leg 1406. More particularly, in at least one embodiment, the staple pocket 1500 can be positioned relative to the staple 1400 such that, as the staple 1400 is ejected from a staple cartridge, for example, the end 1405 of first leg 1404 can enter the first forming cup 1502 and then the end 1407 of second leg 1406 can enter the second forming cup 1504. Further to the above, the end 1405 of first staple leg 1404 can contact the base 1506 of first forming cup 1502 such that the first compressive force F1 can be applied to the first leg 1404 and, similarly, the end 1407 of second staple leg 1406 can contact the base 1508 of second forming cup 1504 such that the second compressive force F2 can be applied to the second leg 1406.

[0236] In various embodiments, further to the above, the first forming cup 1502 can comprise an inside portion 1510 and an outside portion 1512, wherein, when the end 1405 of first staple leg 1404 enters into the first forming cup 1502, the end 1405 can enter into the outside portion 1512. Upon entering into the outside portion 1512 of forming cup 1502, the end 1405 can contact base 1506 and, owing to a concave curve of base 1506, the end 1405 can be directed inwardly toward the inside portion 1510. More particularly, referring now to FIGS. 77-81, the base 1506 can be curved toward tissue-contacting surface 1501 such that, as the staple leg 1404 contacts the base 1506, the end 1405 can be directed downwardly, i.e., away from tissue-contacting surface 1501, and inwardly along the curved concave surface toward an inflection point 1595. In various embodiments, the inflection point 1595 can represent the point in which the concave surface of base 1506 will begin to deflect the end 1405 of first leg 1404 upwardly toward the tissue-contacting surface 1501. In various embodiments, the radius of curvature r of the concave surface can be constant, or at least substantially constant, in the longitudinal direction along the length thereof as illustrated in FIGS. 80 and 81. In certain embodiments, the radius of curvature r of the concave surface of base 1506 can be consistent across the width of base 1506 between a first interior sidewall 1516 and a first exterior sidewall 1517. In any event, as the end 1405 of first leg 1404 is advanced into the inside portion 1510 of forming cup 1502, the end 1405 can come into contact with a radius transition 1514 positioned intermediate the base 1506 and the first interior sidewall 1516. In such embodiments, the radius transition 1514 can be configured to direct the end 1405 against the first interior sidewall 1516.

[0237] As illustrated in FIG. 76, further to the above, the first interior sidewall 1516 can be oriented at an angle with respect to staple pocket longitudinal axis 1599. In certain embodiments, the first interior sidewall 1516 can be oriented at an acute angle, such as 10 degrees, for example, with respect to longitudinal axis 1599. In various embodiments, the first interior sidewall 1516 and the longitudinal axis 1599 may be neither perpendicular nor parallel to one another. In any event, the first interior sidewall 1516 can extend through the axis 1599 such that a first portion of the first interior sidewall 1516 is positioned on a first side 1515 of axis 1599 and a second portion of the first interior sidewall 1516 is positioned on a second side 1517 of axis 1599. In various embodiments, as a result, the first interior sidewall 1516 can extend between the first outside portion 1512 and the first inside portion 1510. When the end 1405 of first leg 1404 contacts the first interior sidewall 1516, as described above, the end 1405 can be directed along the first interior sidewall 1516 away from longitudinal axis 1599 such that the staple leg 1404 is bent away from the common plane of staple 1400 toward the first side 1515 of axis 1599. As the end 1405 of first leg 1404 is directed along, or bent by, the first interior sidewall 1516, as described above, the staple leg 1404 can also be directed, or bent, by base 1506. Stated another way, the first sidewall 1516 and the first base 1506 can co-operate to deform the first staple leg 1404 such that end 1405 is re-directed toward the base 1402 and, at the same time, to a first side of the base 1402 as described above. At some point during the insertion of first staple leg 1404 into first forming cup 1502, the end 1405 of first staple leg 1404 can emerge from the first inside portion 1510 of first forming cup 1502 and, as the staple leg 1404 is further deformed by the staple pocket 1500, the end 1405 can be directed along the first axis 1414 (FIG. 83) as described above.

[0238] In various embodiments, further to the above, the first interior sidewall 1516 can extend along an interior side of the first base 1506, wherein, in at least one embodiment, the first forming cup 1502 can further comprise a first exterior sidewall 1517 extending along an opposite side of the first base 1506. In certain embodiments, similar to the above, the first forming cup 1502 can further comprise a transition radius 1519 positioned intermediate the base 1506 and the
exterior sidewall 1517. In at least one embodiment, referring now to FIG. 76, the exterior sidewall 1517 can extend in a direction which is parallel, or at least substantially parallel, to the staple pocket longitudinal axis 1599. As also illustrated in FIG. 76, the first interior sidewall 1516 and the first exterior sidewall 1517 can extend in directions which are transverse to one another. In at least one embodiment, the interior sidewall 1516 can extend at an acute angle, such as approximately 15 degrees, for example, with respect to the exterior sidewall 1517. In various embodiments, as a result, the outside portion 1512 of first forming cup 1502 can be wider than the inside portion 1510. In at least one such embodiment, the width of the outside portion 1512 and the inside portion 1510 can taper between a first width and a second width.

[0239] In various embodiments, referring once again to FIG. 76, the outside portion 1512 of first forming cup 1502 can comprise a first outside wall 1513 which can extend in a direction which is perpendicular to the first exterior wall 1517 and/or the longitudinal axis 1599 and can define the outermost portion of forming cup 1502. In at least one embodiment, further to the above, the width of the first outside wall 1513 can be such that the outside portion 1512 can capture the end 1405 of first leg 1404 and guide it into the inside portion 1510 of cup 1502 as described above. In at least one such embodiment, the first outside wall 1513 can be at least as wide as the diameter of the first leg 1404. In certain embodiments, the first forming cup 1502 can further comprise a channeling surface 1528 surrounding the first inner portion 1510 and the first outer portion 1512 which can be configured to guide the staple leg 1404 into and/or out of the forming cup 1502. In various embodiments, the inside portion 1510 can further comprise an inside wall 1511 which can define the innermost portion of forming cup 1502. Similar to the above, the inside wall 1511 can also define the narrowest portion of forming cup 1502. In at least one embodiment, the width of the inside wall 1511 may be the same, or at least substantially the same, as the diameter of first leg 1404 such that the inside wall 1511 can control the location in which the end 1405 emerges from staple forming cup 1502.

[0240] In various embodiments, further to the above, the second forming cup 1504 can comprise an inside portion 1520 and an outside portion 1522, wherein, when the end 1407 of second staple leg 1406 enters into the second forming cup 1504, the end 1407 can enter into the outside portion 1522. Upon entering into the outside portion 1522 of forming cup 1504, the end 1407 can contact base 1508 and, owing to a concave curve of base 1508, the end 1407 can be redirected inwardly toward the inside portion 1520. More particularly, similar to the above, the base 1508 can be curved toward tissue-contacting surface 1501 such that, as the staple leg 1406 contacts the base 1508, the end 1407 can be directed downwardly, i.e., away from tissue-contacting surface 1501, and inwardly along the curved concave surface toward an inflection point 1596. In various embodiments, the inflection point 1596 can represent the point in which the concave surface of base 1508 will begin to deflect the end 1407 of second leg 1406 upwardly toward the tissue-contacting surface 1501. In various embodiments, the radius of curvature, r, of the concave surface can be constant, or at least substantially constant, in the longitudinal direction along the length thereof, similar to the base 1506 of first forming cup 1502 illustrated in FIGS. 80 and 81. In any event, as the end 1407 of second leg 1406 is advanced into the inside portion 1520 of forming cup 1504, the end 1407 can come into contact with a radius transition 1524 positioned intermediate the base 1508 and a second interior sidewall 1526. In such embodiments, the radius transition 1524 can be configured to direct the end 1407 against the second interior sidewall 1526.

[0241] As illustrated in FIG. 76, further to the above, the second interior sidewall 1526 can be oriented at an angle with respect to staple pocket longitudinal axis 1599. In certain embodiments, the second interior sidewall 1526 can be oriented at an acute angle, such as 10 degrees, for example, with respect to longitudinal axis 1599. In various embodiments, the second interior sidewall 1526 and the longitudinal axis 1599 may be neither perpendicular nor parallel to one another. In any event, the second interior sidewall 1526 can extend through the axis 1599 such that a first portion of the second interior sidewall 1526 is positioned on a first side 1515 of axis 1599 and a second portion of the second interior sidewall 1526 is positioned on a second side 1517 of axis 1599. In various embodiments, as a result, the second interior sidewall 1526 can extend between the second outside portion 1522 and the inside portion 1520. When the end 1407 of second leg 1406 contacts the interior sidewall 1526, as described above, the end 1407 can be directed along the interior sidewall 1526 such that the staple leg 1406 is bent away from the common plane of staple 1400 toward the second side 1517 of axis 1599. As the end 1407 of second leg 1406 is directed along, and bent by, the interior sidewall 1526, as described above, the staple leg 1406 can also be directed, and bent, by base 1508. Stated another way, the second interior sidewall 1526 and the base 1508 can co-operate to deform the second staple leg 1406 such that end 1407 is re-directed toward the base 1402 and, at the same time, toward a second, or opposite, side of the base 1402 as described above. At some point during the insertion of second staple leg 1406 into second forming cup 1504, the end 1407 of second staple leg 1406 can emerge from the second inside portion 1520 of second forming cup 1504 and, as the staple leg 1406 is further deformed by the staple pocket 1500, the end 1407 can be directed along the second axis 1416 (FIG. 83) as described above.

[0242] In various embodiments, further to the above, the second interior sidewall 1526 can extend along an interior side of the second base 1508, wherein, in at least one embodiment, the second forming cup 1504 can further comprise a second exterior sidewall 1527 extending along an opposite side of the second base 1508. In certain embodiments, similar to the above, the second forming cup 1504 can further comprise a transition radius 1529 positioned intermediate the base 1508 and the exterior sidewall 1527. In at least one embodiment, referring now to FIG. 76, the exterior sidewall 1527 can extend in a direction which is parallel, or at least substantially parallel, to the staple pocket longitudinal axis 1599. As also illustrated in FIG. 76, the second interior sidewall 1526 and the second exterior sidewall 1527 can extend in directions which are transverse to one another. In at least one embodiment, the interior sidewall 1526 can extend at an acute angle, such as approximately 15 degrees, for example, with respect to the exterior sidewall 1527. In various embodiments, as a result, the outside portion 1522 of second forming cup 1504 can be wider than the inside portion 1520. In at least one such embodiment, the width of the outside portion 1522 and the inside portion 1520 can taper between a first width and a second width.

[0243] In various embodiments, referring once again to FIG. 76, the outside portion 1522 of second forming cup 1504
can comprise a second outside wall 1523 which can extend in a direction which is perpendicular to the second exterior wall 1527 and/or the longitudinal axis 1599 and can define the outermost portion of forming cup 1504. In at least one embodiment, further to the above, the width of the second outside wall 1523 can be such that the outside portion 1522 can capture the end 1407 of second leg 1406 and guide it into the inside portion 1520 of cup 1504 as described above. In at least one such embodiment, the second outside wall 1523 can be at least as wide as the diameter of the second leg 1406. In certain embodiments, the second forming cup 1504 can further comprise a channeling surface 1529 surrounding the second inner portion 1520 and the second outer portion 1522 which can be configured to guide the staple leg 1406 into and/or out of the forming cup 1504. In various embodiments, the inside portion 1520 can further comprise an inside wall 1521 which can define the innermost portion of forming cup 1504. Similar to the above, the inside wall 1521 can also define the narrowest portion of forming cup 1504. In at least one embodiment, the width of the inside wall 1521 may be the same, or at least substantially the same, as the diameter of second leg 1406 such that the inside wall 1521 can control the location in which the end 1407 emerges from staple forming cup 1504.

[0244] As discussed above, referring again to FIGS. 76-78, the first forming cup 1502 can comprise a first interior side wall 1516 and the second forming cup 1504 can comprise a second interior side wall 1526. As illustrated in FIG. 76, the first inside portion 1510 of forming cup 1502 can be positioned in close proximity to, or close relation to, the second inside portion 1520 of forming cup 1504 such that the first interior side wall 1516 can be positioned adjacent to the second interior side wall 1526. In at least one embodiment, the first interior portion 1510, or at least a substantial portion thereof, can be offset from the staple pocket longitudinal axis 1599 in the first direction 1515 while the second interior portion 1520, or at least a substantial portion thereof, can be offset from the longitudinal axis 1599 in the second direction 1517. In various embodiments, the staple pocket 1500 can comprise a wall 1530 positioned intermediate the first inside portion 1510 and the second inside portion 1520, wherein a first side of wall 1530 can comprise the first interior side wall 1516 and wherein a second side of wall 1530 can comprise the second interior side wall 1526. In at least one such embodiment, the first interior side wall 1516 can be parallel, or at least substantially parallel, to the longitudinal axis 1599. More particularly, in at least one embodiment, the first interior side wall 1516 can define a first plane and the second interior side wall 1526 can define a second plane, wherein the first plane and the second plane can be parallel, or at least substantially parallel, to one another. In various embodiments, referring again to FIGS. 77 and 78, the first interior side wall 1516 can be perpendicular, or at least substantially perpendicular, to the tissue-contacting surface 1501 and, similarly, the second interior side wall 1526 can be perpendicular, or at least substantially perpendicular, to the tissue-contacting surface 1501.

[0245] In various embodiments, further to the above, the first interior side wall 1516 can comprise a first vertical portion 1516a which is perpendicular, or at least substantially perpendicular, to the tissue-contacting surface 1501. In at least one embodiment, the first vertical portion 1516a can extend through, or transect, the longitudinal axis 1599. In various embodiments, the first vertical portion 1516a can extend along the entirety of, or only a portion of, the first interior side wall 1516. Similarly, the second interior side wall 1526 can comprise a second vertical portion 1526a which is perpendicular, or at least substantially perpendicular, to the tissue-contacting surface 1501. In at least one embodiment, such a second vertical portion 1526a can extend through, or transect, the longitudinal axis 1599. In various embodiments, the second vertical portion 1526a can extend along the entirety of, or only a portion of, the second interior side wall 1526. During the deployment of staple 1400, further to the above, the end 1405 of first leg 1404 can be in contact with the first vertical portion 1516a of first interior side wall 1516 at the same time the end 1407 of second leg 1406 is in contact with the second vertical portion 1526a of second interior side wall 1526. In such circumstances, the first vertical portion 1516a and the second vertical portion 1526a can comprise a vertical trap. More particularly, the vertical portions 1516a and 1526a can co-operate to control, deflect, and bend the staple legs 1404 and 1406 in opposite directions, i.e., in directions to the sides of a common plane, as described above, when the legs 1404 and 1406 come into contact with the interior sidewalls 1516 and 1526 of forming cups 1502 and 1504, respectively. For example, referring again to FIG. 78, the first vertical portion 1516a can be configured to deflect and bend the staple leg 1404 to a first side of base 1402 and the second vertical portion 1526a can be configured to deflect and bend the staple leg 1406 to a second, or opposite, side of base 1402.

[0246] In various embodiments, further to the above, the vertical trap comprising vertical portions 1516a and 1526a can extend along the entire length of the first and second interior side walls 1516 and 1526, while, in other embodiments, the vertical trap may extend along only a portion of the sidewalls 1516 and 1526. In at least one embodiment, the vertical trap can be approximately 0.05 inches long, i.e., the overlap of the first vertical surface 1516a and the second vertical surface 1526a can be approximately 0.05 inches, for example, along the lengths of interior surfaces 1516 and 1526. In various embodiments, the length of the vertical trap can be between approximately 0.03 inches and approximately 0.10 inches, for example. In certain embodiments, the length of the vertical trap can be approximately twice the radius of curvature (r) of the curved concave surface of base 1506, for example. In various embodiments, the length of the vertical trap can be approximately equal to the radius of curvature (r) of base 1506, for example. In at least one embodiment, the length of the vertical trap can be between approximately 0.5r and approximately 2r, for example. In various embodiments, further to the above, the vertical trap can extend through the longitudinal axis 1599 of staple pocket 1500 such that, in at least one embodiment, at least a portion of the vertical trap can be positioned on a first side and/or a second side of axis 1599. In certain embodiments, the vertical trap can extend through the central portions of the first and second forming cups 1502 and 1504.

[0247] In various embodiments, the first interior side wall 1516 can further comprise a first angled portion which, in at least one embodiment, can be oriented at an acute angle with respect to the tissue-contacting surface 1501. In at least one such embodiment, the first angled portion can be positioned outwardly with respect to the first vertical portion 1516a. In certain embodiments, the first interior side wall 1516 can comprise an angled portion positioned toward the outside portion 1512 which can become progressively more perpendicular toward the inside portion 1510 of the first forming cup.
1502 until the angled portion transitions into the first vertical portion 1516a. In various embodiments, the second interior sidewall 1526 can further comprise a second angled portion which, in at least one embodiment, can be oriented at an acute angle with respect to the tissue-contacting surface 1501. In at least one such embodiment, the second angled portion can be positioned outwardly with respect to the second vertical portion 1526a. In certain embodiments, the second interior sidewall 1526 can comprise an angled portion positioned toward the outside portion 1522 which can become progressively more perpendicular toward the inside portion 1520 of the second forming cup 1504 until the angled portion transitions into the second vertical portion 1526a.

[0248] In various embodiments, referring now to FIG. 85A, the staple pocket 1500 can be configured to deform the first staple leg 1404 such that the first end 1405 is deflected at a first distance X1 from baseline 1401. Similarly, the second staple leg 1406 can be deformed such that the second end 1407 is deflected a second distance X2 from baseline 1401. In certain embodiments, the distance X1 and the distance X2 can be the same, or at least substantially the same. In various other embodiments, the distances X1 and X2 can be different. In at least one such embodiment, the first leg 1404 can be deformed such that the first end 1405 is position closer to base 1402 than the second end 1407, for example. In such embodiments, the first axis 1414 of deformed staple leg 1404 and the second axis 1416 of deformed staple leg 1406 may be non-parallel. More particularly, in at least one embodiment, the first axis 1414 can extend at a first angle with respect to baseline 1401 and the second axis 1416 can extend at a second angle with respect to baseline 1401 where the second angle is different than the first angle. In various embodiments, the first leg 1404 and the second leg 1406 can extend across midline 1403 at different angles. In certain other embodiments, the first leg 1404 and the second leg 1406 can be extend at different angles with respect to baseline 1401 although one or both of the legs 1404 and 1406 may not extend across the midline 1403.

[0249] In various embodiments, further to the above, a surgical stapler can comprise a staple pocket which can be configured to deform one staple leg of staple 1400 such that it lies within, or substantially within, a common plane with base 1402 and, in addition, deform the other staple leg of staple 1400 to a side of base 1402 as described above. In at least one embodiment, the first leg 1404 can be deformed such that it extends through midline 1403 in a direction which is coplanar, or at least substantially co-planar, with base 1402 and, in addition, the second leg 1406 can be deformed such that it extends through midline 1403 in a direction which is transverse to the plane. Stated another way, in at least one embodiment, axis 1414 and baseline 1401 of staple 1400 can be coplanar, or at least nearly co-planar, with one another while second axis 1416 can extend in a direction which extends through such a plane. In certain embodiments, at least one of the first leg 1404 and the second leg 1406 may not extend through the midline 1403.

[0250] In various embodiments, further to the above, the staple pocket 1500 can be configured to deform the staple legs 1404 and 1406 of staple 1400 simultaneously, or at least substantially simultaneously. In at least one embodiment, the base 1506 of first forming cup 1502 can contact end 1405 of first staple leg 1404 at the same time, or at least substantially the same time, that the base 1508 of second forming cup 1504 contacts end 1407 of second staple leg 1406. In certain other embodiments, a staple pocket can be configured to deform the staple legs 1404 and 1406 sequentially. In at least one such embodiment, a first forming cup can be brought into contact with the first staple leg 1404 before a second forming cup is brought into contact with the second staple leg 1406, for example. In various other embodiments, although not illustrated, a surgical staple can comprise more than two staple legs, such as three staple legs or four staple legs, for example, and a staple pocket can comprise a corresponding quantity of staple forming cups for deforming the staple legs.

[0251] In various embodiments, further to the above, the wire comprising the surgical staple 1400 can comprise a circular, or at least substantially circular, cross-section. In various other embodiments, referring now to FIGS. 86-89, a surgical staple, such as staple 1400, for example, can comprise a non-circular cross-section. In at least one embodiment, the staple 1600 can comprise a base 1602, a first leg 1604, and a second leg 1606, wherein the base 1602 and legs 1604 and 1606 can be comprised of a continuous wire. In various embodiments, the continuous wire can comprise a rectangular cross-section, for example. In at least one embodiment, referring to FIG. 89, the rectangular cross-section can comprise a base (b) and a height (h), wherein the base (b) can be defined relative to a central lateral axis (x), and wherein the height (h) can be defined relative to a central longitudinal axis (y). In various circumstances, the rectangular cross-section can be defined as having two moments of inertia, i.e., a first moment of inertia (lx) defined with respect to axis (x) and a second moment of inertia (ly) defined with respect to axis (y). In at least one circumstance, the first moment of inertia (lx) can be calculated as (b*h^3)/12 while the second moment of inertia (ly) can be calculated as (b^3*h)/12. Although staple 1600 comprises a rectangular, or at least substantially rectangular cross-section, any other suitable non-circular cross-section can be utilized, such as oblate, elliptical, and/or trapezoidal cross-sections, for example.

[0252] As illustrated in FIG. 89, the base (b) of surgical staple 1600 is larger than the height (h) and, in view of the above, the moment of inertia (ly) of the rectangular cross-section is larger than the moment of inertia (lx). In various embodiments, as a result, the moment of inertia ratio, i.e., ly/lx, of the rectangular cross-section can be greater than 1.0. In certain embodiments, the moment of inertia ratio can be between approximately 2.0 and approximately 2.7, for example. In certain other embodiments, the moment of inertia ratio can be between approximately 1.1 and approximately 3.0, for example. As a result of the above, the leg 1604 is more likely to bend about axis (x) than about axis (y) when a force such as compressive load F1, for example, is applied to the leg 1604. In any event, absent all other considerations, the leg 1604, in such embodiments, is more likely to bend within a common plane defined by the staple 1600 when it is in its undeformed state than bend to a side of staple base 1602. In various embodiments, however, a surgical stapler comprising an anvil and staple pocket in accordance with the embodiments described herein, such as staple pocket 1500, for example, can be utilized to cause the legs 1604 and 1606 of staple 1600 to bend out of their common plane when they are deformed. In such embodiments, this lateral deflection can occur despite the fact that the moment of inertia ly, which resists such twisting, is greater than the moment of inertia lx. As illustrated in FIG. 88, the first leg 1604 of staple 1600 can be deformed such that it is bent relative to both axis (x) and axis (y) of its cross-section and, as a result, the first staple leg
1604 can be twisted or deformed such that the end 1605 of first staple leg 1604 is positioned on a first side of base 1602. Similarly, the second leg 1606 can be deformed such that it is bent relative to both axis (x) and axis (y) of its cross-section and, as a result, the second staple leg 1606 can be twisted or deformed such that the end 1607 of second staple leg 1606 is positioned on a second side of base 1602.

[0253] In various embodiments, referring now to FIG. 90, a surgical staple, such as surgical staple 1700, for example, can comprise a base 1702 and, in addition, a first leg 1704 and a second leg 1706 extending from base 1702. In certain embodiments, similar to the above, the base 1702, the first leg 1704, and the second leg 1706 can lie, or at least substantially lie, in a common plane when the staple 1700 is an undeformed, or undeployed, configuration, i.e., a configuration prior to being deformed by an anvil of a surgical stapler, for example. In the deformed or deployed configuration of staple 1700, as illustrated in FIG. 90, the first leg 1704 can be deformed such that end 1705 points toward base 1702 and second leg 1706. More particularly, in at least one embodiment, the end 1705 can lie along, or with respect to, a first axis 1714 which is oriented at angle with respect to midline 1703. Similarly, the second leg 1706 can be deformed such that end 1707 points toward base 1702 and first leg 1704. More particularly, in at least one embodiment, the end 1707 can lie along, or with respect to, a second axis 1716 which is oriented at angle with respect to midline 1703. In various embodiments, the ends 1705 and 1707 of legs 1704 and 1706 may not cross mid-line 1703. In certain embodiments, similar to the above, the end 1705 of first leg 1704 may be deformed such that it extends to a first side of base 1702 and the end 1707 of second leg 1706 may be deformed such that it extends to a second, or opposite, side of base 1702 such that legs 1704 and 1706 are not entirely positioned in-plane with base 1702 in their deformed configuration, for example.

[0254] In various embodiments, a surgical staple, such as staple 1800 (FIG. 91), for example, can comprise a base 1802, a first leg 1804, and a second leg 1806, wherein the staple 1800 can comprise a substantially U-shaped configuration in its undeformed, or undeployed, configuration. In at least one such embodiment, legs 1804 and 1806 can extend in a perpendicular, or at least substantially perpendicular, direction with respect to base 1802. In various circumstances, the staple 1800 can be deformed into a b-shaped configuration as illustrated in FIG. 91. In at least one such embodiment, the first leg 1804 can be bent downwardly toward base 1802 such that axis 1814 extending through end 1805 is perpendicular, or at least substantially perpendicular, to baseline 1801. Similarly, the second leg 1806 can be bent downwardly toward base 1802 such that axis 1816 extending through end 1807 is perpendicular, or at least substantially perpendicular, to baseline 1801. In at least one such circumstance, the legs 1804 and 1806 can be bent such that axes 1814 and 1816 are parallel, or at least substantially parallel, to one another. In various embodiments, referring again to FIG. 91, the staple legs 1804 and 1806 can be deformed such that they do not cross central line 1803. The staple legs 1804 and 1806 can be deformed such that they remain in-plane, or at least substantially in-plane, with base 1802.

[0255] Various examples described below are envisioned which incorporate one or more aspects of the various embodiments described above. Such examples are exemplary and various aspects of various embodiments described in this application can be combined in a single embodiment. In each of the examples described below, the surgical staple can comprise a base defining a baseline, a first leg and a second leg which extend from the base, and a midline midway between the first leg and the second leg.

Example 1

A surgical staple can be deformed such that:

<table>
<thead>
<tr>
<th>First Leg</th>
<th>Second Leg</th>
</tr>
</thead>
<tbody>
<tr>
<td>Crosses the midline (FIG. 83)</td>
<td>Crosses the midline (FIG. 83)</td>
</tr>
<tr>
<td>Extends in-plane, or substantially in-plane, with the base (FIG. 91)</td>
<td>Extends out of plane with the base (FIG. 85)</td>
</tr>
<tr>
<td>The end extends in a non-perpendicular direction with the baseline (FIG. 83)</td>
<td>The end extends in a non-perpendicular direction with the baseline (FIG. 83)</td>
</tr>
</tbody>
</table>

Example 2

A surgical staple can be deformed such that:

<table>
<thead>
<tr>
<th>First Leg</th>
<th>Second Leg</th>
</tr>
</thead>
<tbody>
<tr>
<td>Crosses the midline (FIG. 83)</td>
<td>Crosses the midline (FIG. 83)</td>
</tr>
<tr>
<td>Extends out of plane with the base (FIG. 85) to the same side of the base as the second leg, the distance X1 being different than X2 (FIG. 85A)</td>
<td>Extends out of plane with the base (FIG. 85) to the same side of the base as the first leg, the distance X1 being different than X2 (FIG. 85A)</td>
</tr>
<tr>
<td>The end extends in a non-perpendicular direction with the baseline (FIG. 83)</td>
<td>The end extends in a non-perpendicular direction with the baseline (FIG. 83)</td>
</tr>
</tbody>
</table>
Example 3
[0258]  A surgical staple can be deformed such that:

<table>
<thead>
<tr>
<th>First Leg</th>
<th>Second Leg</th>
</tr>
</thead>
<tbody>
<tr>
<td>Does not cross the midline (FIG. 90)</td>
<td>Does not cross the midline (FIG. 90)</td>
</tr>
<tr>
<td>Extends out of plane with the base (FIG. 85) to a first side of the base,</td>
<td>Extends out of plane with the base (FIG. 85) to a second side of the base,</td>
</tr>
<tr>
<td>the distance X1 being different than X2 (FIG. 85A)</td>
<td>the distance X1 being different than X2 (FIG. 85A)</td>
</tr>
<tr>
<td>The end extends in a non-perpendicular direction with the baseline (FIG.</td>
<td>The end extends in a non-perpendicular direction with the baseline (FIG. 83)</td>
</tr>
<tr>
<td>83)</td>
<td></td>
</tr>
</tbody>
</table>

Example 4
[0259]  A surgical staple can be deformed such that:

<table>
<thead>
<tr>
<th>First Leg</th>
<th>Second Leg</th>
</tr>
</thead>
<tbody>
<tr>
<td>Does not cross the midline (FIG. 90)</td>
<td>Does not cross the midline (FIG. 90)</td>
</tr>
<tr>
<td>Extends out of plane with the base (FIG. 85) to the same side of the base</td>
<td>Extends out of plane with the base (FIG. 85) to the same side of the base,</td>
</tr>
<tr>
<td>as the second leg, the distance X1 being different than X2 (FIG. 85A)</td>
<td>the distance X1 being different than X2 (FIG. 85A)</td>
</tr>
<tr>
<td>The end extends in a non-perpendicular direction with the baseline (FIG.</td>
<td>The end extends in a non-perpendicular direction with the baseline (FIG. 83)</td>
</tr>
<tr>
<td>83)</td>
<td></td>
</tr>
</tbody>
</table>

Example 5
[0260]  A surgical staple can be deformed such that:

<table>
<thead>
<tr>
<th>First Leg</th>
<th>Second Leg</th>
</tr>
</thead>
<tbody>
<tr>
<td>Does not cross the midline (FIG. 90)</td>
<td>Does not cross the midline (FIG. 90)</td>
</tr>
<tr>
<td>Extends in-plane, or substantially in-plane, with the base (FIG. 91)</td>
<td>Extends out of plane with the base (FIG. 85)</td>
</tr>
<tr>
<td>The end extends in a perpendicular direction with the baseline (FIG. 91)</td>
<td>The end extends in a non-perpendicular direction with the baseline (FIG. 83)</td>
</tr>
</tbody>
</table>

Example 6
[0261]  A surgical staple can be deformed such that:

<table>
<thead>
<tr>
<th>First Leg</th>
<th>Second Leg</th>
</tr>
</thead>
<tbody>
<tr>
<td>Crosses the midline (FIG. 83)</td>
<td>Does not cross the midline (FIG. 90)</td>
</tr>
<tr>
<td>Extends out of plane with the base (FIG. 85) to a first side of the base,</td>
<td>Extends out of plane with the base (FIG. 85) to a second side of the base,</td>
</tr>
<tr>
<td>the distance X1 being different than X2 (FIG. 85A)</td>
<td>the distance X1 being different than X2 (FIG. 52A)</td>
</tr>
<tr>
<td>The end extends in a non-perpendicular direction with the baseline (FIG.</td>
<td>The end extends in a non-perpendicular direction with the baseline (FIG. 83)</td>
</tr>
<tr>
<td>83)</td>
<td></td>
</tr>
</tbody>
</table>
Example 7

[0262] A surgical staple can be deformed such that:

<table>
<thead>
<tr>
<th>First Leg</th>
<th>Second Leg</th>
</tr>
</thead>
<tbody>
<tr>
<td>Crosses the midline (FIG. 83)</td>
<td>Does not cross the midline (FIG. 90)</td>
</tr>
<tr>
<td>Extends out of plane with the base (FIG. 85) to the same side of the base as the second leg, the distance X₁ being different than X₂ (FIG. 85A)</td>
<td>Extends out of plane with the base (FIG. 85) to the same side of the base as the second leg, the distance X₁ being different than X₂ (FIG. 85A)</td>
</tr>
<tr>
<td>The end extends in a non-perpendicular direction with the baseline (FIG. 83)</td>
<td>The end extends in a non-perpendicular direction with the baseline (FIG. 83)</td>
</tr>
</tbody>
</table>

Example 8

[0263] A surgical staple can be deformed such that:

<table>
<thead>
<tr>
<th>First Leg</th>
<th>Second Leg</th>
</tr>
</thead>
<tbody>
<tr>
<td>Crosses the midline (FIG. 83)</td>
<td>Does not cross the midline (FIG. 90)</td>
</tr>
<tr>
<td>Extends out of plane with the base (FIG. 85)</td>
<td>Extends in plane, or substantially in-plane, with the base (FIG. 91)</td>
</tr>
<tr>
<td>The end extends in a non-perpendicular direction with the baseline (FIG. 83)</td>
<td>The end extends in a perpendicular direction to the baseline (FIG. 91)</td>
</tr>
</tbody>
</table>

Example 9

[0264] A surgical staple can be deformed such that:

<table>
<thead>
<tr>
<th>First Leg</th>
<th>Second Leg</th>
</tr>
</thead>
<tbody>
<tr>
<td>Crosses the midline (FIG. 83)</td>
<td>Does not cross the midline (FIG. 90)</td>
</tr>
<tr>
<td>Extends in-plane, or substantially in-plane, with the base (FIG. 91)</td>
<td>Extends out of plane with the base (FIG. 85)</td>
</tr>
<tr>
<td>The end extends in a non-perpendicular direction with the baseline (FIG. 83)</td>
<td>The end extends in a non-perpendicular direction with the baseline (FIG. 83)</td>
</tr>
</tbody>
</table>

[0265] Several of the deformed staples described above comprise one or more staple legs which cross the mid-line of the staple base. In various embodiments, as a result, the deformed staple legs may at least partially overlap with one another. More particularly, the deformed staple legs, when viewed from the side, may co-operate to traverse the staple base from one end to the other leaving no gap therebetween. Such embodiments can be particularly useful, especially when used to staple vascular tissue. More specifically, the overlapping staple legs can compress blood vessels within the tissue regardless of where the blood vessels extend through the staple. Staples having gaps between the legs, or legs which do not extend along the entire length of the staple base when deformed, may not be able to properly compress every blood vessel in the tissue and, as a result, one or more blood vessels may leak.

[0266] In various embodiments, further to the above, a surgical instrument can be configured to deploy a plurality of staples 1600 in the manner described above and illustrated in FIG. 88. In at least one such embodiment, similar to the above, the surgical stapler can deploy the staples 1600 in a sequential manner along a staple path and/or in a simultaneous manner, for example. In various embodiments, a surgical instrument can be configured to deploy a plurality of staples 1600 in the manner described above and illustrated in FIG. 88. In at least one such embodiment, similar to the above, the surgical stapler can deploy the staples 1600 in a sequential manner along a staple path and/or in a simultaneous manner, for example. In various embodiments, further to the above, a surgical instrument can be configured to deploy a plurality of staples 1700 in the manner described above and illustrated in FIG. 90. In at least one such embodiment, the surgical stapler can deploy the staples 1700 in a sequential manner along a staple path and/or in a simultaneous manner, for example.

[0267] In various embodiments, referring now to FIGS. 103-108, a surgical stapling instrument 2100 can comprise, similar to the above, a first housing portion 2102 and a second housing portion 2104 which can be operably connected to one another by a latch 2180. Latch 2180 can comprise a frame 2184 which can be pivotally mounted to a frame 2110 of first housing portion 2102. In use, the latch 2180 can be configured to engage a frame 2114 of second housing portion 2104 and draw the second housing portion 2104 toward the first housing portion 2102 and move the anvil support portion 2130 of second housing portion 2104 toward the staple cartridge support portion 2124 of first housing portion 2102. In various
embodiments, the first housing portion 2102, the second housing portion 2104, and the latch 2180 can each comprise one or more contoured outer housings or gripping portions, for example. In at least one such embodiment, the first housing portion 2102 can comprise an outer housing 2108, the second housing portion 2104 can comprise an outer housing 2112, and the latch 2180 can comprise an outer housing 2186. The surgical stapling instrument 2100 can further comprise a firing actuator 2204 which can, similar to the above, be selectively positioned on opposite sides of the surgical stapling instrument. More particularly, further to the above, the actuator 2204 can be selectively positioned on a first side of the housing portions 2102, 2104 such that the actuator 2204 can be moved distally along the first side or selectively positioned on a second side of the housing portions 2102, 2104 such that the actuator 2204 can be moved distally along the second side. In at least one embodiment, the first housing portion 2102 and the second housing portion 2104 can define one or more slots 2118 therebetween which can permit the actuator 2204 to be moved along the first and second sides. In at least one such embodiment, the slots 2118 can be connected by an intermediate slot 2331 which can extend around and/or through the proximal end of the surgical stapling instrument 2100.

[0269] Further to the above, referring to FIGS. 103-111, the firing actuator 2204 can be rotatably mounted to a drive bar 2220, wherein, in at least one embodiment, the actuator 2204 can be rotatably mounted to the drive bar 2220 via a connecting link 2206. Referring primarily to FIGS. 109-111, the actuator 2204 can be rotated between an intermediate, or neutral, position (FIG. 109) in which the drive bar 2220 is locked in position and cannot be advanced distally and an unlocked position (FIG. 110) in which the drive bar 2220 and the actuator 2204 are ready to be fired distally. Although FIG. 110 illustrates the actuator 2204 in an unlocked position on the first side of the surgical stapling instrument housing, the actuator 2204 can also be moved into an unlocked position on the opposite, or second, side of the surgical stapling instrument housing. The following example, although discussed in connection with the actuator 2204 being moved along the first side of the housing, is also applicable in connection with the actuator 2204 being moved along the second side of the housing. In any event, the actuator 2204 and the first housing portion 2102, for example, can comprise various interlocking features which can prevent, or at least limit, relative movement between the drive bar 2220 and the first housing portion 2102. More particularly, in at least one embodiment, the first housing portion 2102 can comprise one or more slots and/or one or more projections which can cooperate with one or more slots and/or one or more projections of the actuator 2204 such that the drive bar 2220 cannot be advanced distally until the actuator 2204 has been sufficiently rotated out of its neutral position and into an unlocked position. In various embodiments, the proximal end of the first housing portion 2102 can comprise an end post 2107 which can include a retention slot 2213 configured to receive at least a portion of actuator link 2206, such as retention projection 2214, therein. When the actuator 2204 is in its neutral position, the retention projection 2214 is positioned in the retention slot 2213 and neither the actuator 2204 nor the drive bar 2220 can be advanced distally. As the actuator 2204 is rotated toward an unlocked position, the retention projection 2214 can move out of the retention slot 2213 in end post 2107 and into a receiving slot 2215 in drive bar 2220 as illustrated in FIG. 110. In at least one embodiment, the drive bar 2220 and actuator 2204 can remain in a locked condition until the retention projection 2214 has completely exited the retention slot 2213. Thereafter, the actuator 2204 can be advanced distally. In addition to or lieu of the above, the end post 2107 can further comprise a retention wall 2211 which can, similar to the above, impede the distal movement of actuator 2204 and drive bar 2220. More particularly, the actuator link 2206 can further comprise a retention projection 2216 which can be positioned behind, or distally with respect to, the retention wall 2211 when the actuator 2204 is in its neutral position and, owing to such alignment, the retention wall 2211 can provide a bearing surface preventing the distal movement of retention projection 2216. Once actuator 2204 has been sufficiently rotated out of its neutral position toward an unlocked position, the retention projection 2216 can be moved to a position in channel 2217 which is out of longitudinal alignment with the retention wall 2211 thereby permitting relative longitudinal movement therebetween.

[0270] As described above, once the actuator 2204 has been moved into an unlocked position (FIG. 110), the actuator 2204 can be advanced distally into a fired position (FIG. 111). In such circumstances, referring now to FIG. 106, a force can be applied to the actuator 2204 in order to advance drive bar 2220 distally and incise tissue and/or deploy staples from a staple cartridge as described above. In such circumstances, the force can rotate and seat the actuator 2204 in a fully-deployed, or at least nearly fully-deployed, position. In at least one embodiment, the drive bar 2220 can comprise one or more stops, such as stops 2221 (FIG. 111), for example, which can limit the rotation of the actuator 2204 in the distal direction. In at least one such embodiment, the drive bar 2220 can comprise a first stop 2221 configured to limit the rotation of the actuator 2204 toward the first side of the instrument and a second stop 2221 configured to limit the rotation of the actuator toward the second side of the instrument. In certain embodiments, referring to FIG. 106, the stops 2221 can be configured such that the actuator 2204 is positioned along an axis which is perpendicular, or at least substantially perpendicular, to a longitudinal axis 2299 of the surgical stapling instrument 2100. Such a position of actuator 2204 is also illustrated in FIG. 104. Referring now to FIG. 107, a force can be applied to the actuator 2204 in order to retract the actuator 2204 proximally. In such circumstances, the force can cause the actuator 2204 to rotate proximally until it comes into contact with the first housing portion 2102 and/or the second housing portion 2104. Such a position of actuator 2204 is also illustrated in FIG. 105 wherein the actuator 2204 can be positioned against lock rail 2131 and/or lock rail 2132, for example, in order to prevent any further rotation of the actuator 2204.

[0270] In various embodiments, as described above, the latch 2180 can be utilized to lock the first housing portion 2102 and the second housing portion 2104 together. In certain embodiments, the actuator 2204 can be utilized to limit the relative movement between the housing portions 2102, 2104 and/or move the housing portions 2102, 2104 toward one another. In at least one embodiment, referring primarily to FIGS. 103, 104, and 108, the actuator 2204 can comprise a recess, or channel, 2130 which can be configured to receive the lock rails 2131 and 2132 when the actuator 2204 is moved along the first side of the surgical stapling instrument 2100 or, alternatively, receive the lock rails 2133 and 2134 when the actuator 2204 is moved along the second side of the surgical stapling instrument 2100. In either event, the recess 2130 can
be configured to capture an opposing set of rails, such as rails 2131 and 2132, for example, and prevent, or at least limit, relative movement therebetween. More particularly, in at least one embodiment, the recess 2130 can comprise a first bearing surface 2135 positioned opposite the first lock rail 2131 and a second bearing surface 2136 positioned opposite the second lock rail 2132 such that the bearing surfaces 2135, 2136 can prevent, or at least limit, the movement of the first housing portion 2102 and the second housing portion 2104 away from one another. In some circumstances, gaps may exist between the bearing surfaces 2135, 2136 and the lock rails 2131, 2132, respectively, when the bearing surfaces 2135, 2136 are adjacent to the lock rails 2131, 2132 while, in other circumstances, the bearing surface 2135 may contact the lock rail 2131 and/or the bearing surface 2136 may contact the lock rail 2132, for example. In use, in various circumstances, the actuator 2204 can be moved from its neutral position (FIG. 109) into an unlocked position (FIG. 110) wherein, in such a position, the recess 2130 can be aligned with either a set of rails 2131, 2132 or a set of rails 2133, 2134 depending on whether the actuator 2204 has been rotated to the first or second side. When the actuator 2204 is advanced distally, in some circumstances, the actuator 2204 may contact the rails and cam, or drive, the rails toward each other. In such circumstances, the first housing portion 2102 and the second housing portion 2104 can be cannulated, or driven, toward one another.

In various embodiments, as described above, the actuator 2204 can receive, capture, and/or engage a lock rail extending from each of the first housing portion 2102 and the second housing portion 2104. In various alternative embodiments, the actuator 2204 can be configured to receive, capture, and/or engage two or more lock rails extending from the first housing portion 2102 and/or the second housing portion 2104. In certain embodiments, the first housing portion 2102 and/or the second housing portion 2104 can comprise one or more lock channels which can be configured to receive at least a portion of the actuator. In various embodiments, the housing portions and the actuator of the surgical stapling instrument can comprise any suitable lock portions which can be configured to receive, align, retain, capture, lock, move, cam, and/or limit the movement of the surgical instrument housing portions. In various embodiments, referring primarily to FIGS. 106 and 107, the lock rails 2131, 2132, 2133, and/or 2134 can extend longitudinally along the stapling instrument 2100 such that, in at least one embodiment, they extend in a longitudinal direction from the proximal end of the surgical stapling instrument 2100 toward the distal end of the instrument which is parallel, or at least substantially parallel, to longitudinal axis 2299. Furthermore, in at least one embodiment, the lock rails 2131, 2132, 2133, and/or 2134 can extend in directions which are parallel, or at least substantially parallel, to one another.

In various embodiments, further to the above, a surgical staple can be comprised of titanium, such as titanium wire, for example. In certain embodiments, a surgical staple can be comprised of an alloy comprising titanium, aluminum, and/or vanadium, for example. In at least one embodiment, the surgical staple can be comprised of surgical stainless steel and/or an alloy comprised of cobalt and chromium, for example. In any event, the surgical staple can be comprised of metal, such as titanium, and a metal oxide outer surface, such as titanium oxide, for example. In various embodiments, the metal oxide outer surface can be coated with a material. In certain embodiments, the coating material can be comprised of polytetrafluoroethylene (PTFE), such as Teflon®, and/or a tetrafluoroethylene (TFE) such as ethylene-tetrafluoroethylene (ETFE), perfluoralkoxyethylene-tetrafluoroethylene (PEA), and/or fluorinated Ethylene Propylene (FEP), for example. Certain coatings can comprise silicon. In various embodiments, such coating materials can prevent, or at least inhibit, further oxidation of the metal. In certain embodiments, the coating materials can provide one or more lubricious surfaces against which the anvil, or staple pockets, can contact the staples in order to reduce the friction force therebetween. In various circumstances, lower friction forces between the staples and the staple pockets can reduce the force required to deform the staples.

The devices disclosed herein can be designed to be disposed of after a single use, or they can be designed to be used multiple times. In either case, however, the device can be reconditioned for reuse after at least one use. Reconditioning can include any combination of the steps of disassembly of the device, followed by cleaning or replacement of particular pieces, and subsequent reassembly. In particular, the device can be disassembled, and any number of the particular pieces or parts of the device can be selectively replaced or removed in any combination. Upon cleaning and/or replacement of particular parts, the device can be reassembled for use either at a reconditioning facility, or by a surgical team immediately prior to a surgical procedure. Those skilled in the art will appreciate that reconditioning of a device can utilize a variety of techniques for disassembly, cleaning/replacement, and reassembly. Use of such techniques, and the resulting reconditioned device, are all within the scope of the present application.

Preferably, the invention described herein will be processed before surgery. First, a new or used instrument is obtained and if necessary cleaned. The instrument can then be sterilized. In one sterilization technique, the instrument is placed in a closed and sealed container, such as a plastic or TYVEK bag. The container and instrument are then placed in a field of radiation that can penetrate the container, such as gamma radiation, x-rays, or high-energy electrons. The radiation kills bacteria on the instrument and in the container. The sterilized instrument can then be stored in the sterile container. The sealed container keeps the instrument sterile until it is opened in the medical facility.

While this invention has been described as having exemplary designs, the present invention may be further modified within the spirit and scope of the disclosure. This application is therefore intended to cover any variations, uses, or adaptations of the invention using its general principles. Further, this application is intended to cover such departures from the present disclosure as come within known or customary practice in the art to which this invention pertains.

What is claimed is:

1. A surgical stapling instrument, comprising:
   a first side;
   a second side;
   a longitudinal axis;
   a first housing member, comprising:
   a staple cartridge attachment portion configured to operably support a staple cartridge; and
   a first lock rail extending along said longitudinal axis;
   a second housing member, comprising:
   a jaw member configured to operably support an anvil configured to deform staples deployed from the staple cartridge; and
   a second lock rail extending along said longitudinal axis;
   a pusher bar configured to move a staple driver relative to said staple cartridge attachment portion and said jaw member; and
an actuator configured to receive a force thereto, wherein said actuator is configured to be moved between a first position and a second position, wherein said actuator is configured to be moved along said first side when said actuator is in said first position, wherein said actuator is configured to be moved along said second side when said actuator is in said second position, and wherein said actuator further comprises at least one lock rail receiver configured to receive said first lock rail and said second lock rail.

2. The surgical stapling instrument of claim 1, wherein said at least one lock rail receiver comprises a first bearing surface configured to engage said first lock rail and a second bearing surface configured to engage said second lock rail, and wherein said first bearing surface and said second bearing surface are configured to move said first housing member and said second housing member toward each other.

3. The surgical stapling instrument of claim 1, wherein said at least one lock rail receiver comprises a first bearing surface positioned adjacent to said first lock rail and a second bearing surface positioned adjacent to said second lock rail, and wherein said first bearing surface and said second bearing surface are configured to limit relative movement between said first lock rail and said second lock rail.

4. The surgical stapling instrument of claim 3, wherein said first bearing surface and said second bearing surface define opposite sides of a channel.

5. The surgical stapling instrument of claim 1, wherein said first lock rail and said second lock rail extend from said first side, wherein said at least one lock rail receiver is configured to receive said first lock rail and said second lock rail when said actuator is in said first position, wherein said first housing member further comprises a third lock rail extending from said second side, wherein said second housing member further comprises a fourth lock rail extending from said second side, and wherein said at least one lock rail receiver is configured to receive said third lock rail and said fourth lock rail when said actuator is in said second position.

6. The surgical stapling instrument of claim 1, further comprising said anvil.

7. The surgical stapling instrument of claim 6, further comprising said staple cartridge.

8. A surgical stapling instrument, comprising:
   a first side;
   a second side;
   a first housing member, comprising:
   a first proximal end portion;
   a first distal end portion;
   a staple cartridge attachment portion configured to operably support a staple cartridge; and
   a first longitudinal lock portion extending from said first proximal end portion toward said first distal end portion;
   a second housing member, comprising:
   a second proximal end portion;
   a second distal end portion;
   a jaw member configured to operably support an anvil configured to deform staples deployed from the staple cartridge; and
   a second longitudinal lock portion extending from said second proximal end portion toward said second distal end portion;
   a drive member configured to move a staple driver relative to said staple cartridge attachment portion and said jaw member; and
   an actuator configured to receive a force thereto, wherein said actuator is configured to be moved between a first position and a second position, wherein said actuator is configured to be moved along said first side when said actuator is in said first position, wherein said actuator is configured to be moved along said second side when said actuator is in said second position, and wherein said actuator further comprises at least one lock rail receiver configured to receive said first lock rail and said second lock rail.

9. The surgical stapling instrument of claim 8, wherein said at least one actuator lock portion comprises a first bearing surface configured to engage said first longitudinal lock portion and a second bearing surface configured to engage said second longitudinal lock portion, and wherein said first bearing surface and said second bearing surface are configured to move said first housing member and said second housing member toward each other.

10. The surgical stapling instrument of claim 8, wherein said at least one actuator lock portion comprises a first bearing surface positioned adjacent to said first longitudinal lock portion and a second bearing surface positioned adjacent to said second longitudinal lock portion, and wherein said first bearing surface and said second bearing surface are configured to limit relative movement between said first housing member and said second housing member.

11. The surgical stapling instrument of claim 10, wherein said first bearing surface and said second bearing surface define opposite sides of a channel.

12. The surgical stapling instrument of claim 8, wherein said first longitudinal lock portion and said second longitudinal lock portion extend from said first side, wherein said at least one actuator lock portion is configured to receive said first longitudinal lock portion and said second longitudinal lock portion when said actuator is in said first position, wherein said first housing member further comprises a third longitudinal lock portion extending from said second side, wherein said second housing member further comprises a fourth longitudinal lock portion extending from said second side, and wherein said at least one actuator lock rail portion is further configured to be aligned with said third longitudinal lock portion and said fourth longitudinal lock portion when said actuator is in said second position.

13. The surgical stapling instrument of claim 12, further comprising said staple cartridge.

14. The surgical stapling instrument of claim 8, further comprising said anvil.

15. A surgical stapling instrument, comprising:
   a first side;
   a second side;
   a longitudinal axis;
   a first housing member, comprising:
   a staple cartridge attachment portion configured to operably support a staple cartridge; and
   a first lock rail extending along said longitudinal axis; and
   a second housing member, comprising:
   a jaw member configured to operably support an anvil configured to deform staples deployed from the staple cartridge; and
   a second lock rail extending along said longitudinal axis; a pusher bar configured to move a staple driver relative to said staple cartridge attachment portion and said jaw member; and
an actuator configured to receive a force thereto, wherein said actuator is configured to be moved between a first position and a second position, wherein said actuator is configured to be moved along said first side when said actuator is in said first position, wherein said actuator is configured to be moved along said second side when said actuator is in said second position, and wherein said actuator further comprises capturing means for capturing said first lock rail and said second lock rail.

16. The surgical stapling instrument of claim 15, wherein said capturing means comprises a first bearing surface configured to engage said first lock rail and a second bearing surface configured to engage said second lock rail, and wherein said first bearing surface and said second bearing surface are configured to limit relative movement between said first lock rail and said second lock rail.

17. The surgical stapling instrument of claim 15, wherein said capturing means comprises a first bearing surface positioned adjacent to said first lock rail and a second bearing surface positioned adjacent to said second lock rail, and wherein said first bearing surface and said second bearing surface are configured to limit relative movement between said first lock rail and said second lock rail.

18. The surgical stapling instrument of claim 17, wherein said first bearing surface and said second bearing surface define opposite sides of a channel.

19. The surgical stapling instrument of claim 15, wherein said first lock rail and said second lock rail extend from said first side, wherein said capturing means is configured to receive said first lock rail and said second lock rail when said actuator is in said first position, wherein said first housing member further comprises a third lock rail extending from said second side, wherein said second housing member further comprises a fourth lock rail extending from said second side, and wherein said capturing means is configured to receive said third lock rail and said fourth lock rail when said actuator is in said second position.

20. The surgical stapling instrument of claim 15, further comprising said staple cartridge.

21. The surgical stapling instrument of claim 15, further comprising said anvil.

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